

(57) Abstract
 An essentially purified preparation of human Lung Cancer-Associated Protein (LCA), a monoclonal antibody specific for LCA, and a method of detecting LCA in a biological sample, which method includes the steps of (1) contacting the sample with an antibody containing the monoclonal antibody specific for LCA, and (2) detecting immunogenic complex formation between the antibody and a constituent of the biological sample, such immune complex formation being indicative of the presence of LCA in the biological sample.

(54) Title: LUNG CANCER-ASSOCIATED PROTEIN

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35 including (1) a first reagent including a first
 Also within the invention is an immunoassay kit
 the hybridioma DF-L1.

30 carcinoma of the lung; one such Mab is that produced by
 of patients with either adenocarcinoma or squamous cell
 23 units of LCAp/ml) with serum samples from at least 70%
 elevated levels (i.e., above the normal cuttoff level of
 no. ATCC HTB 55), and (2) detects circuлатing antigen at
 CALU-3 cells (American Type Culture Collection accession
 LCAp antigen purified from the supernatant of cultured
 25 as an antibody which (1) forms an immune complex with
 monoclonal antibody specific for LCAp is herein defined
 of immune complex formation in the control sample. A
 complex formation in the biological sample to the amount
 monoclonal antibody; and comparing the amount of immune
 20 control sample with a second aliquot containing the
 containing a known amount of LCAp); contacting the
 purified or in a mixture, such as a serum
 sample containing a standard amount of LCAp (either
 include the additional steps of providing a control
 15 biological sample. The method of the invention may
 being indicative of the presence of LCAp in the
 (for example, by ELISA), such immune complex formation
 the antibody and a constituent of the biological sample between
 for LCAp, and detecting immune complex formation between
 10 sample containing a monoclonal antibody (Mab) specific
 contacting the biological sample with an aliquot or
 another mammal), which method includes the steps of
 mucosal scrapings, or biopsied tissue from a human or
 biological sample (e.g., blood, serum, urine, sputum,
 5 invention provides a method for detecting LCAp in a
 incorporated by reference). The immunoassay of the
 al., Cancer Research 51:3838-3842, 1991, herein
 higher than those for normal individuals (Mammals et
 serum of lung cancer patients at levels significantly

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35 E. coli and other enteric bacteria) ! Salmonella heat-
bacteria) ; the so-called "Shiga-like" toxins (produced by
Shiga toxin (produced by various strains of Shigella
cholera toxin (produced by Vibrio cholerae bacteria) ;
toxins such as abrin, modeccin, volkenstein, and viscumin;
30 toxins! Pseudomonas exotoxin A; ricin and other plant
into the immune system of the invention include diphteria
occurring proteaceous toxins that could be incorporated
portion by a peptide bond. Examples of naturally-
in which the antibody portion is linked to the toxin
recombinant DNA molecule would result in an immune toxin
antibody as a single polypeptide: expression of this
both the toxin and an LCA-P-binding portion of the
genetically engineering a hybrid DNA molecule encoding
methodology, or, if the toxin is a protein, by means of
such conjugation may be accomplished by known chemical
binding fragment thereof, conjugated to a toxin molecule.
e.g., the MAb produced by hybridoma DF-L1, or an LCA-P-
immune toxin is an LCA-P-specific monoclonal antibody
immune toxin in which the antibody portion of the
In another aspect, the invention includes an
the antibody from the medium.

25 recombinant DNA molecule would result in an immune toxin
antibody as a single polypeptide: expression of this
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the invention includes an
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35 disclosures set forth herein.

In yet another aspect, the invention features a vaccine for immunizing a human against tumors which express LCA, or an antiigenic fragment thereof, in a portion of LCA, or an antiidiotype antibody such as Freud's, pharmaceuticality-acceptable carrier. The vaccine would enhance the recipient's immune response to the differentially also include an adjuvant such as Freud's, to one of ordinary skill in the art, given the vaccine. Formulation of such a vaccine would be routine to enhance the recipient's immune response to the vaccine.

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In yet another aspect, the invention features a vaccine for immunizing a human against tumors which express LCA, or an antiidiotype antibody such as Freud's, to enhance the immune response to the vaccine.

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In yet another aspect, the invention features a LCA.

Invasive determination of the presence, location, or absence of an LCA-expressing tumor in animal, which would be particularly useful for monitoring the condition of a patient being treated for a tumor known to express LCA.

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Invasive determination of the presence, location, or absence of an LCA-expressing tumor in animal, which bound to a given site being indicative of a tumor at that lung tissue) of the animal, a high level of such label presence of the detectable label bound to a tissue (e.g., detecting (e.g., by radioimaging, using scintigraphy) the tumor; introducing the imaging agent into the animal; and the steps of identifying an animal suspended off having a useful for detecting tumors *in situ* by a method including technique, or *in vivo*). Such an imaging agent would be such as a radionuclide (for example, ¹²⁵I, ¹³¹I,

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also within the invention is an imaging agent in which an LCA-specific monoclonal antibody, or an LCA-binding fragment thereof, is linked to a detectable label which an LCA-specific monoclonal antibody, or an LCA-

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also within the invention is an imaging agent in cells which express LCA on their surfaces.

Invention would be useful for targeting and killing tumor nucleides such as yttrium. An immuno-toxin of the emitting radiionuclides such as astatine and β -emitting anticancer agents such as doxorubicin, as well as a non-protective toxins include known cytotoxic labile enterotoxin; and E. coli heat-labile enterotoxin.

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Non-protective toxins include known cytotoxic nucleides such as yttrium. An immuno-toxin of the emitting radiionuclides such as astatine and β -emitting anticancer agents such as doxorubicin, as well as a-

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SUBSTITUTE FIGURES

- Fig. 1 is a set of graphs illustrating the results of indirect immunofluorescence of human lung carcinoma cells with Mabs DF-L1 and DF-L2. Suspensions of CALU-3, SK-MES, A-549, and ZR-75-1 cells were incubated with Mab DF-L1 (heavy solid line), Mab DF-L2 (dotted line), or an isotype-identical control antibody (thin solid line). After a second incubation with fluorescein-conjugated goat anti-mouse IgG1, the cells were analyzed by flow cytometry.
- Fig. 2 is an indirect analysis of human tumor cells with Mab DF-L1. A, human lung carcinomas; B, other human tumors. Extracells of the indicated cells were subjected to 3-10% gradient SDS-PAGE. The proteins were then transferred to nitrocellulose paper and monitored for reactivity with Mab DF-L1 and 125 I-labeled sheep anti-mouse immunoperoxidase staining of primary human lung carcinomas and normal lung tissue. Formalin-fixed tissue sections were stained with Mab DF-L1 using an avidin-biotin-peroxidase method. A, adenocarcinoma; B, squamous cell carcinoma; C, normal alveolar lining cells; D, normal bronchus (arrow, terminal web of brush border).
- Fig. 3 is a set of photographs showing mouse immunoglobulin. Mabs DF-L1 and 125 I-labeled sheep anti-tissue sections were stained with Mab DF-L1 using an avidin-biotin-peroxidase method. A, immunoperoxidase staining of primary human lung carcinomas and normal lung tissue. Formalin-fixed tissue sections were stained with Mab DF-L1 using an avidin-biotin-peroxidase method. B, immunoperoxidase staining of primary human lung carcinomas and normal lung tissue. Formalin-fixed tissue sections were stained with Mab DF-L1 using an avidin-biotin-peroxidase method. C, normal alveolar lining cells; D, squamous cell carcinoma; E, normal alveolar lining cells; F, normal bronchus (arrow, terminal web of brush border).
- Fig. 4 is an autoradiogram showing immunoprecipitation of [3 H]proline-labeled CALU-3 cells. CALU-3 cells were labeled with [3 H]proline for 48 h. Cells were subjected to immunoprecipitation with Mab DF-L1, or an IgG1 control Mab. The immunoprecipitate was analyzed by 3-15% SDS-PAGE and autoradiography. Rd, M_r in thousands.
- Fig. 5 is an immunoblot illustrating the effects of tunicamycin on DF-L1 antigen. CALU-3 cells were grown in tunicamycin on DF-L1 antigen. Fig. 5 shows the effects of tunicamycin on DF-L1 antigen. CALU-3 cells were grown in tunicamycin on DF-L1 antigen. Fig. 5 shows the effects of tunicamycin on DF-L1 antigen.

Drawings

The drawings are first briefly described.

Detailed Description

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lung carcinoma.

NSCLC, non-small cell lung carcinoma; SCLC, small cell levels. A normal cut-off value of 23 units/ml was used. and patients with lung cancer were assayed for LCAP 30 with lung cancer. Plasma samples from normal subjects Fig. 10 is a graph showing LCAP levels in patients plotted as a histogram.

subjects with LCAP levels within the indicated ranges was from normal subjects was assayed, and the number of 25 distribution of LCAP levels in normal subjects. Plasma Fig. 9 is a histogram illustrating the bars, SD; ABS, absorbances.

patients with metastatic lung cancer. Points, mean; normal subjects; closed circles and squares denote 20 a function of dilution. Open circles and squares denote patients with lung cancer were assayed for LCAP levels as on LCAP levels. Plasma samples from normal subjects and Fig. 8 is a graph showing the effect of dilution bars, SD.

closed circles, Day 2; ■, Day 3; ▲, Day 4. Points, mean; determined for each calibrator. Open circles, Day 1; consecutive days. Absorbances (ABS) at 490 nm were 15 LCAP calibrator curves. LCAP calibrators containing 0, Fig. 7 is a graph showing the reproducibility of 3 antigen. kd, M⁻¹ in thousands.

analyses with MAb DF-L1. Lane 9, 1 µg of purified CALU- carcinomas (Lanes 5-8) were subjected to immunoblot 20, 100, and 200 units/ml were assayed in duplicate on a LCAP calibrator curves. LCAP calibrators containing 0, Fig. 6 is an immunoblot analysis of human plasma 5 samples analyzed with MAb DF-L1. Plasma specimens (3 µl) subjected to immunoblot analysis with MAb DF-L1. kd, M⁻¹ in the presence of 10 µg/ml tunicamycin for 24 h and then in the presence of 10 µg/ml tunicamycin for 24 h and then 10 Fig. 6 is an immunoblot analysis of human plasma 3 antigen. kd, M⁻¹ in thousands.

subjected to immunoblot analysis with MAb DF-L1. kd, M⁻¹ 15 samples analyzed with MAb DF-L1. Plasma specimens (3 µl) from normal subjects (Lanes 1-4) and patients with lung carcinomas (Lanes 5-8) were subjected to immunoblot 20, 100, and 200 units/ml were assayed in duplicate on a LCAP calibrator curves. LCAP calibrators containing 0, Fig. 7 is a graph showing the reproducibility of 3 antigen. kd, M⁻¹ in thousands.

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patients with metastatic lung cancer. Points, mean; normal subjects; closed circles and squares denote 25 a function of dilution. Open circles and squares denote patients with lung cancer were assayed for LCAP levels as on LCAP levels. Plasma samples from normal subjects and Fig. 10 is a graph showing LCAP levels in patients plotted as a histogram.

subjects with LCAP levels within the indicated ranges was from normal subjects was assayed, and the number of 30 distribution of LCAP levels in normal subjects. Plasma Fig. 11 is a graph showing LCAP levels in patients plotted as a histogram.

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- Fig. 11 is a set of graphs showing serial LCAP levels in patients with lung cancer. Serial LCAP levels were determined and compared with clinical course during effective treatment in patients with lung cancer. Of 3 patients treated for lung cancer were determined and adenoacarcinoma; C, small cell carcinoma. Patients, mean; bars, SD.
- Fig. 12 is a set of graphs showing LCAP spikes during effective treatment in patients with lung cancer. Serial LCAP levels from a patient with Stage IIIA adenocarcinoma of the lung (12A) and from two patients with limited stage small cell carcinoma of the lung (12B, C) were monitored during chemotherapy and radiotherapy. No evidence of detectable disease. Dotted line = 23 U/ml.
- Fig. 13 is a graph showing serial LCAP levels after complete resection of non-small cell carcinoma of the lung. Serial LCAP levels were monitored daily from nine patients with stage I or II non-small cell carcinoma of the lung. Of the lung following complete surgical resection of non-lung malignancies. Dotted line = 23 U/ml.
- Fig. 14 is a graph of LCAP levels in patients with tumor. Horizontal dotted line = 23 U/ml.
- Fig. 15 is a graph showing serial LCAP levels with non-lung malignancies. Dotted line = 23 U/ml.
- Fig. 16 is a graph illustrating a typical case of lung malignancies. Dotted line = 23 U/ml.
- The invention, including points plotted on the curve for the invention, the invention curve obtained with the immunoassay kit of calibration curve following a graph illustrating a typical case of lung malignancies. Dotted line = 23 U/ml.
- Immunoassay of mice with an extract of a primary human lung adenocarcinoma resulted in a panel of two hypothetical test samples.
- Human lung adenocarcinoma closely related by subjecting a lung cancer cell line closely-related, high molecular weight glycoproteins, was extracted from a number of lung and breast carcinoma cell lines. LCAP antigen, which is actually a group of hybrigrams designated DF-L1 and DF-L2, which react with monoclonal antibodies, including those produced by the human lung adenocarcinoma cells, in a panel of
- 35 first identified by subjecting a lung cancer cell line closely-related, high molecular weight glycoproteins, was extracted from a number of lung and breast carcinoma cell lines. LCAP antigen, which is actually a group of hybrigrams designated DF-L1 and DF-L2, which react with human lung adenocarcinoma cells, in a panel of

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Mab Production. Mabs were generated by techniques analogous to those described previously (Kufe et al.,
Hybridoma 3:223-232, 1984). Briefly, BALB/c mice were immunized with an extract of a primary human adenocarcinoma of the lung. Mouse spleen cells were fused with P3X63-Ag8.653 myeloma cells (ATCC CRL 1580) and hybridomas cloned three times by limiting dilution in Dulbecco's modified Eagle's medium with 4.5 g/liter glucose, 10% fetal bovine serum, 10% NCTC-109 medium supplement (Sigma), 1% sodium pyruvate, 1% nonessential amino acids, 200 mM L-glutamine, 1% tylosin (Sigma Chemical Co.), and 1% penicillin/streptomycin. Hybridoma cells were injected into pristane-primed BALB/c mice and the Mabs purified from ascites using a Protein A-Sepharose column (Bioprod, Richmond, VA). Yields ranged from 0.5-9.0 mg purified antibody/ml ascites fluid.

Culture of Human Tumor Cells. The human lung adenocarcinoma cell line CALU-3 (ATCC HTB 55), the squamous cell carcinoma SK-MES (ATCC HTB 58), and the undifferentiated lung carcinoma A-549 (ATCC CCL 185) were cultured in T25 flasks at 37°C in 5% CO₂.

Immunofluorescence Assay. Cells were seeded onto chamber slides and allowed to grow until confluent. Cells were fixed with 4% paraformaldehyde for 15 min at room temperature. After fixation, cells were permeabilized with 0.2% Triton X-100 for 10 min. Cells were then incubated with primary antibodies overnight at 4°C. Secondary antibodies were applied for 1 h at room temperature. Cells were viewed under a fluorescence microscope (Olympus IX71) equipped with a color camera (QImaging QICQ-1394) and analyzed using Image Pro Plus software (Media Cybernetics).

Flow Cytometry. Cells were harvested and washed with PBS. Cells were stained with primary antibodies for 1 h at room temperature. Cells were washed and stained with secondary antibodies for 1 h at room temperature. Cells were analyzed using a flow cytometer (BD FACSCanto II) and FlowJo software (Tree Star).

Western Blotting. Cells were lysed with RIPA buffer and protein concentration was determined using a bicinchoninic acid (BCA) assay. Proteins were separated on 10% SDS-PAGE gels and transferred to polyvinylidene difluoride (PVDF) membranes. Membranes were probed with primary antibodies overnight at 4°C. Secondary antibodies were applied for 1 h at room temperature. Membranes were visualized using an ECL kit (GE Healthcare).

RT-PCR. Total RNA was extracted from cells using Trizol reagent. cDNA was synthesized using a reverse transcriptase kit (Thermo Fisher Scientific). PCR was performed using primers specific for target genes. PCR products were resolved on 1.5% agarose gels and visualized using ethidium bromide staining.

Statistical Analysis. Statistical significance was determined using Student's t-test or one-way ANOVA. P < 0.05 was considered statistically significant.

Materials and Methods

I. DETECTION AND CHARACTERIZATION OF LCAP

extract to SDS-PAGE and immunoblotting the gel with the monoclonal antibody DF-L1. An immunoassay utilizing this antibody was subsequently used to detect LCAp in serum samples from human patients, leading to the discovery that LCAp is present in significantly higher levels in samples from patients with lung cancer or certain other types of cancer, than in those from normal, healthy individuals. These studies are described in detail below. Also described below is an immunoassay kit for the detection and quantitation of circulating LCAp, useful for evaluating and monitoring patients with lung cancer, and potentially other types of cancer as well.

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35 Malvern, PA). The supernatant of the cell extract was rabbit anti-mouse immunoglobulin (Organon-Tekniska, protein A-conjugated Sepharose CL-4B (Pharmacia) with immunoprecipitation (IP) complex was formed by incubating centrifuged at 10,000 x g for 10 min. An

30 fluoride/20 mM phenanthroline) and the cell extract was protease inhibitors (5 mM EDTA/20 mM phenylmethylenol 1 mM Tris-HCl lysate buffer (pH 8.0) in the presence of Amersham). The cells were lysed with a 1% Nonidet-40/50 medium containing 150 μ Ci [3 H]glucosamine (40 Ci/mmol; 25 Amersham). CALU-3 cells were also incubated in complete fresh medium with 150 μ Ci [3 H]proline (130 Ci/mmol; 6 h. The cells were then incubated for 48-72 h with in proline-free medium supplemented with dialyzed FBS for immunoprecipitation. CALU-3 cells were incubated 20 for cytoplasmic and membrane-staining patte

25 ren. Reactivity was assessed on a 0-3+ visual scale μ g/ml. Primary antibody was used at a concentration of 0.1 CA). Vectastain ABC Kit, Vectored Laboratories, Burlingame, using an avidin-biotin-peroxidase staining technique 15 formalin-fixed paraffin-embedded tissue were stained for immunoperoxidase staining. Four-mm sections of washed, dried, and exposed with Kodak X-OMAT film.

15 (Amersham, Arlington Heights, IL). The blot was then with 125 I-labeled sheep anti-mouse immunoglobulin incubated with primary antibody, washed, and incubated nitrocellulose membrane was blocked with 5% BSA, transfer (14). Following Western transfer, the Coomassie blue or PAS(13) or subjected to Western 3-10% gradient SDS-PAGE(12). Gels were then stained with immunoblotting. Antigen samples were analyzed by sorter (Coulter, Hialeah, FL).

5 Analyzed on a dual-beam fluorescence-activated cell azide. The cells were again washed extensively and maintained) for 1 h at 40°C in the presence of 0.2% sodium

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human tumor cell lines in an indirect ELISA. Although determined by reactivity against extracts of various study. The specificity of these Mabs was first designated DF-T1 and DF-T2, were chosen for further panel of monoclonal antibodies. Two IgG1 Mabs, 30 human lung adenocarcinoma resulted in the production of a Immunoisolation of BALB/c mice with an extract of a primary REACTIVITY WITH HUMAN TUMOR CELLS LINES.

RESULTS

25 ANALYSIS.

20 Separation of IgG1 Mabs from the culture supernatant was performed by Coomassie and PAs staining or immunoblot followed by SDS-PAGE approximately 50%. Purity was monitored by SDS-PAGE hypothesized, and stored at -20°C. Antigen yield was MgCl₂, extensively dialyzed against water, concentrated, Sepharose CL-4B (Pharmacia). Antigen was eluted with 3 M covalently coupling 4 mg of Mab/ml of CNBr-activated and applied to a Mab affinity column prepared by fractions with antigen activity were collected, pooled, and monitored by absorbance at 280 nm and by ELISA. 15 4B (Pharmacia) sizing column. Fractions were collected concentrated medium was then applied to a Sepharose CL- (Amicon, Danvers, MA) on a YM30 membrane. The approximately 30 times in a stirred ultrafiltration cell x g for 20 min to remove debris, and concentrated spent medium was collected, pooled, centrifuged at 10,000 cells. Cells were grown for 3 days past confluence, and antigen was purified from spent culture medium of CALU-3 (Amicon, Danvers, MA) on a YM30 membrane. The approximately 30 times in a stirred ultrafiltration cell x g for 20 min to remove debris, and concentrated spent medium was collected, pooled, centrifuged at 10,000 10 cells. Cells were grown for 3 days past confluence, and antigen was purified from spent culture medium of CALU-3 antigen Purification. High molecular weight monitored by fluorography.

5 Subjected to SDS-PAGE under reducing conditions, and then incubated overnight at 4°C with Mab followed by irrelevant mouse IgG1 Mab. The precleared extract was pre-cleared by incubating with the IP complex and an irrelevant mouse IgG1 Mab. The precleared extract was then incubated for 1 h with IP complex, washed extensively, then incubated overnight at 4°C with Mab followed by

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the spectrum of reactivity with these antibodies was nearly identical, there were certain quantitative differences. For example, both reacted with all of the lung and breast carcinoma cells (Table 1). In contrast, MAb DF-L1 reacted with an extract of OVCAR ovarian carcinoma cells, while there was no detectable reactivity against these cells using MAb DF-L2 (Table 1). Neither MAb reacted with cell lines derived from human leukemias or melanoma (Table 1).

Indirect immunofluorescence was similarly used to determine whether the epitopes recognized by these antibodies are expressed on the surface of lung carcinoma cell lines. As compared to a control antibody, both MAb DF-L1 and DF-L2 demonstrated binding to the CALU-3, SK-MES, and A-549 lung carcinoma cell lines (Fig. 1). However, the patterns of reactivity of the two MAbs against the same cell lines were distinct (Fig. 1).

Similar results were obtained with the ZR-75-1 breast carcinoma cells (Fig. 1). Taken together with the ELISA data, these findings suggested that MAb DF-L1 and MAb DF-L2 react with distinct epitopes.

Immunoblot analysis of extracts from the lung carcinoma cells revealed reactivity with MAb DF-L1 but not with MAb DF-L2. A heterogeneous antigen with an apparent Mr of 350,000-420,000 was detected in CALU-3 cells (Fig. 2A). High molecular weight antigens were similarly detected in SK-MES, CALU-1, and A-549 cell lines (Fig. 2A). Although these reactive species were more homogeneous in size and ranged slightly higher in lines, although these reactive species were more

In contrast with the findings by ELISA, there was no detectable reactivity of this MAb with extracts of the lung and breast carcinomas. In fact, MAb DF-L1 also reacted with a heterogeneous group of high molecular weight antigens in the breast and ovarian carcinoma cell lines (Fig. 2B). In contrast with the ELISA, there was no detectable reactivity of this MAb with extracts of the lung and breast carcinomas. In fact, MAb DF-L1 also reacted with a heterogeneous group of high molecular weight antigens in the breast and ovarian carcinoma cell lines (Fig. 2B).

35 937 or HL-60 leukemia lines (Fig. 2B).

Reactivity with Human Tissues. Formalin-fixed paraffin-embedded sections of tumor and normal tissue were evaluated for reactivity with Mabs DF-L1 and DF-L2 using an immunoperoxidase-staining technique. While there was no detectable staining with Mab DF-L2, certain lung cancer specimens reacted with Mab DF-L1 (Table 2). For example, all adenocarcinomas stained intensely with Mab DF-L1 within the cytoplasm and on apical borders. Mab DF-L1 in the adenocarcinoma sections ranged from 40-100%. Squamous cell carcinomas of the lung expressed the DF-L1 epitope to a lesser degree than adenocarcinomas and DF-L1 epitope to a lesser degree than adenocarcinomas and 100%. Squamous cell carcinomas of the lung expressed the DF-L1 in the adenocarcinoma sections ranged from 40-100%. The percentage of tumor cells reactive with DF-L1 was observed in the central fields, with the more rare positive cells. However, a "pavementing" pattern was observed in the central fields, with the more differentiated areas of the squamous cell cancers had the staining pattern was distinct. Peripherally, poorly the staining pattern was distinct. Peripherally, poorly differentiated areas of the squamous cell cancers had differentiated squamous cells exhibiting intense, rare positive cells. However, a "pavementing" pattern was observed in the central fields, with the more differentiated areas of the squamous cell cancers had the staining pattern was distinct. Peripherally, poorly differentiated areas of the squamous cell cancers had the staining pattern was distinct. Peripherally, poorly differentiated squamous cells exhibiting intense, differentiated squamous cells exhibiting intense, differentiates in mixed histological areas were positive for the epitope. In contrast, sections from five small cell the epithelium. In contrast, sections from five small cell tumors had no detectable Mab DF-L1 reactivity.

Adenocarcinomas of the breast and ovary expressed tumors had no detectable Mab DF-L1 reactivity.

the DF-L1 epitope to varying degrees (Table 3). In contrast, melanomas, sarcomas, and lymphomas had no detectable staining. Several normal tissues reacted with Mab DF-L1 including kidney and lung. In the kidney, only detectable staining could be seen in normal lung, normal alveolar lining cells adjacent to tumor tissue stained for this antigen but primarily with an apical pattern (Fig. 3C). Certain areas of the bronchus also were positive for this antigen but primarily with a localized pattern primarily at the terminal web of the brush border of the lung. While the glomeruli were negative. In the lung, normal the distal collecting ducts reacted with the antigen, while the alveolar lining cells adjacent to tumor tissue stained for this antigen but primarily with an apical pattern (Fig. 3C).

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35 (Table 4). Similar findings were obtained by double-

30 agents was associated with loss of MAb DF-L2 binding

agents was associated with loss of MAb DF-L1 reactivity, exposure of the antigen to these

alkaline/borohydride and periodate had little effect on

for both antibodies (Table 4). In contrast, while

trypsin, treatment was associated with loss of reactivity

determined by dot immunoblotting. Pronase, but not

structure. Reactivity of Mabs DF-L1 and DF-L2 was

with various agents that alter carbohydrate or protein

The purified antigen was subjected to treatment

25 shown).

20 detectable by Coomassie blue and PAs staining (data not

detectable containing proteins or carbohydrate rates were

purified antigen was analyzed by gel electrophoresis. No

then further purified by MAb DF-L1 immunofinity. The

antigen was fractionated on a Sepharose CL-4B column and

from the culture supernatant of CALU-3 cells. The

these antibodies was performed using antigen purified

characterization of the epitopes recognized by

DF-L2 (data not shown).

15 immunoprecipitation was performed with MAb DF-L1 or MAb

by MAb DF-L1 regardless of whether the

analysis with MAb DF-L1. Similar bands were identified

The immunoprecipitates were subjected to immunoblot

immunoprecipitated with either MAb DF-L1 or MAb DF-L2.

experiments, unlabeled CALU-3 cell extracts were

both antibodies (Fig. 4). Moreover, in other

high molecular weight proteins (M_r 350,000-420,000) for

CALU-3 cells labeled with [3 H]proline revealed with same

antigen. Indeed, immunoprecipitation of antigens from

5 suggested that these antibodies recognize the same

patterns of reactivity with Mabs DF-L1 and DF-L2

identification of reactive epitopes. The similar

slight staining (Fig. 3D).

cells. Occasional basal cells also showed

S U B J E C T

(SPP) and N-succinimidyl-4-(maleimido-
reagents, N-succinimidyl-3-(2-pyridyldithio)propionate
40:219-223, 1981) using the two heterobifunctional
92:1413-1418, 1982; Pain et al., J. Immunol. Methods
of published methods (Yoshitake et al., J. Bioc hem.
30 described above was conjugated to HRP by a modification
described previously. Purified MAb prepared as
peroxidase conjugation. Purified MAb prepared as

Materials and Methods

FOR CIRCULATING CAP

II. DEVELOPMENT AND CHARACTERIZATION OF AN IMMUNOASSAY

25 patients (Fig. 6).

to three reactive species were detectable in certain
mobility of the antigen varied among individuals and up
patients with lung cancer (Fig. 6). The electrophoretic
DF-L1 was clearly greater in plasma samples from four
individuals (Fig. 6). In contrast, reactivity with MAb
of antigen were present in samples from four normal
high molecular weight antigen. Low but detectable levels
were monitored by immunoblotting for the presence of this
from normal individuals and patients with lung cancer
15 detection of circulating antigen. Plasma samples
at least in part, of peptide and O-linked carbohydrate.
peptide structure, while the DF-L2 epitope is compacted,
suggested that the DF-L1 epitope residues primarily in a
of the antigen (Fig. 5). Taken together, these findings
effect on antibody reactivity or electrophoretic mobility
glycosylation. However, this agent had no detectable
grown in the presence of tunicamycin to inhibit N-linked
inhibitory effects (Table 4). CALU-3 cells were also
5 abolished antibody binding and trypsin had only partial
MAb DF-L2 (Table 4). Moreover, pronase completely
borohydride treatments predominantly decreased that for
binding of either MAb, while peridate and alkaline-

determinant ELISA. Neuraminidase had little effect on
- 16 -

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SUGGESTED USE OF CALIBRATORS

- 35 and diluted to make arbitrary-defined calibrator LCAp as determined by Western blot analysis, was pooled adenocarcinoma cells, containing high concentrations of spent tissue culture medium from CALU-3 human lung
- 30 Preparation of LCAp calibrators comprehensive analysis.
- 25 greatest extent and was thus chosen for a more distinguishing cancer patients from normal controls to the individual samples (Table 5). The combination DF-L1/DF-L1-HRP plasma samples from lung cancer patients and from normal optimal circulating LCAp detection with a small panel of antibodies were evaluated for culturing cells supernatants were then evaluated for immunoassay combinations that detected soluble LCAp from LCAp levels in solid-phase ELISAs. The five "sandwich" Mabs were evaluated in various combinations to detect reliability to microtiter plates, retaining activity. These reactivities with purified LCAp derived from CALU-3 cells. As determined by direct immunoassay, 3 of the 10 Mabs reactedivity with purified LCAp generated on the basis of a panel of 10 Mabs was generated on the basis of
- 20 reliability to HRP. Moreover, 4 of the Mabs were found to adsorb to HRP. Retained reactivity with purified LCAp after conjugation remained relatively direct immunosay, 3 of the 10 Mabs as determined by direct immunoassay, 3 of the 10 Mabs reactedivity with purified LCAp derived from CALU-3 cells.
- 15 Development of LCAp assay
- 20 Results
- 10 stored at -70°C.
- centrifugation at 100 x g for 15 min, aliquoted, and contained disodium EDTA. Plasma was separated by Review Board. Samples were collected in evacuated tubes according to protocols approved by the International patients at the Dana-Farber Cancer Institute, Boston, MA, Springfield, MO. Plasma samples were collected from greater Ozarks Blood and Tissue Services Blood Bank in subjects were obtained from the American Red Cross plasma samples. Plasma samples from normal subjects were obtained by a point-to-point linear interpolation of the calibration curve.
- 5 samples were determined by a point-to-point linear

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9.52% for the calibrators and 4.13 to 7.61% for the samples (Table 7). Thus, the reproducibility of the assay was satisfactory.

Antigen Recovery. Known quantities of LCAP were added to four plasma samples containing baseline levels of endogenous LCAP. These samples were then assayed, and the expected value and multiplying by 100. Recovery ranged from 96.4 to 106.0% of added LCAP (Table 8).

Interference. Interference studies. The ability of the assay to quantitate LCAP in the presence of potential interferences was investigated.

Intralipid, circulating substances was investigated. Known quantities of LCAP were added to plasma specimens with baseline levels of endogenous LCAP that contained different levels of bilirubin (up to 22.1 mg/dl) These substances are frequently elevated in rheumatoid factor (1:160), or triglycerides (up to 1026 mg/dl) These substances are frequently elevated in plasma specimens and often by nonimmunochemical mechanisms, in the performance of some immunoassays. The specimens were then reassayed in the LCAP assay, and the percentages of recovery was calculated as above (antigen percentage of recovery was calculated as above (antigen recovery)). The data (Table 9) indicated little if any recovery). The data (Table 9) indicated little if any recovery. The data (Table 9) indicated little if any recovery.

Freeze/Thaw. To assess the stability of LCAP in quantitative interference by these substances in the assay. Samples that had been frozen and thawed, specimens were cycled between freezing (-20°C) and thawing (18°-25°C) 9 times and assayed along with aliquots of these same samples that had only been frozen and thawed once. No samples that had only been frozen and thawed once. No freeze-thaw cycles compared with one cycle, as all sample recoveries were >90% of control.

Matched Serum-Plasma Correlation. Serum and EDTA plasma samples were obtained from 156 volunteer donors and assayed for LCAP levels. The values obtained for the plasma samples were regressed on the serum values. The plasma samples were regressed on the serum values. The

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levels were elevated in patients with each histological cancer patients and those from normal controls was highly (Fig. 10). The difference in LCAP levels from lung 30 units/ml and a range of 8 units/ml to >1000 units/ml in 27 of 35 (77.1%) lung cancer patients with a mean of 127 units/ml (Fig. 10). Moreover, LCAP levels were elevated in 27 of 35 (77.1%) lung carcinoma, and small squamous cell carcinoma, large cell carcinoma, 35 and small histological types of lung cancer (adenocarcinoma, circuating LCAP levels. Patients with all four major metastatic lung cancer was screened to determine A panel of 35 plasmas from patients with patients

25

distribution of circulating LCAP levels in lung cancer significant (Mann-Whitney test, $P = 0.01$). LCAP levels in the two groups was statistically overlapped. Nonetheless, the small difference between smokers (0 to 43 units/ml) and nonsmokers (0 to 34 units/ml), the ranges of LCAP levels for smokers (0 to 7 units/ml), was chosen as a reference cut-off value. Although the mean LCAP level for smokers was slightly higher than for nonsmokers (10 units/ml versus 15 units/ml), was chosen as a reference cut-off value.

20

units/ml. A level of the mean plus 2 SDs, or 23 units/ml. Only 13.8% of the samples had levels above 15 units/ml to a high of 43 units/ml. Thirty-five percent of the samples had LCAP levels of 2 units/ml or less, 20 units of 5 units/ml. The levels ranged from a low of 0 value of 5 units/ml. The mean was 7 units/ml (SD \pm 8) with a median of the normals was 7 units/ml (SD \pm 9). The mean value were evaluated for LCAP levels (Fig. 9). The mean value plasma samples from 341 normal blood bank donors

15

10

5

subjects

distribution of circulating LCAP levels in Normal subjects (see following section) was 94%. As a reference cut-off (slope, 0.979. The overall concordance using 23 units/ml as a reference cut-off (see following section) was 94%. Correlation coefficient obtained was 0.966, and the

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- 35 Forty-six percent of patients with chronic obstructive was screened for the determination of LCAP levels.
- A small panel of patients with benign lung disease
- Benign Samples**
- 30 Prostate cancer. None of the patients with gastric patients with ovarian cancer, and 20% of patients with cancer, 50% of patients with breast cancer, 33% of determine the specificity of this assay. Elevated levels
- 25 cancers other than lung cancer were also screened to plasma samples from a small group of patients with
- Non-Lung Cancer Samples**
- 20 Of LCAP prior to chemotherapy rapidly decreased to normal clinical response to therapy (Fig. 11C). Elevated levels from a patient with small cell carcinoma also paralleled progressed, LCAP levels increased. Serial levels of LCAP therapy, LCAP levels decreased, and conversely as disease 6-mo period (Fig. 11B). As the patient responded to
- 15 clinically-documented response to therapy over an initial patient, also with adenocarcinoma, correlated with disease progression. Serial LCAP levels from a second continued for 5 mo prior to clinical documentation of (Fig. 11A); however, his LCAP levels increased
- 10 considered by clinical criteria to have stable disease. For example, one patient with adenocarcinoma was with lung cancer during treatment for metastatic disease.
- LCAP levels were monitored in selected patients
- 5 **Serial LCAP Levels**
- and small cell carcinoma, 4 of 6 (66.7%).
- undifferentiated non-small cell carcinoma, 3 of 3 (100%)!
- squamous cell carcinoma, 4 of 7 (57.1%); other
- type of lung cancer: adenocarcinoma, 16 of 19 (84.2%)!

THE USE OF THE

Sample collection and Clinical Evaluation. Plasma samples collected in EDTA-treated tubes or serum samples were obtained from patients with lung cancer, other malignancies, and a variety of benign disorders. Samples from patients with malignancies were collected from patients at the Dana-Farber Cancer Institute and were stored at -70°C until assayed. These samples were collected within 70°C until assayed. Protocols approved by the Human Studies Committee. Samples from patients with benign disorders. Samples at Sinai Hospital (Detroit) and at Asturias General Hospital (Oviedo, Spain).

Clinical information was obtained by review of the patients' charts without knowledge of LCA P levels. Only patients with histologically-documented lung cancer were eligible. Patients with a prior history of a non-pulmonary malignancy, except basal cell carcinoma of the skin or *in situ* carcinoma of the cervix, were excluded.

For the correlation of serum levels with clinical presentation was available with >30 days between sample collections. Courses, patients were included if more than one sample was available with >30 days between sample collections. For the correlation of serum levels with clinical presentation was available with >30 days between sample collections.

Malignant lesions documented by histologic, clinical, or radiographic evidence of any new progression was defined as the appearance of any new malignant lesions. Patients with >30 days between sample collections were included if more than one sample was available with >30 days between sample collections.

30

Methods and Materials

5 pulmonary disease (COPD) and 54% of patients with pneumonia had elevated LCAP levels. One patient with tuberculosi s had a normal LCAP level. Despite the relatively high percentage of patients with benign disease that had elevated LCAP levels, the means of the LCAP levels were 23.8 and 23.9 units/ml for the COPD and pneumonia patients, respectively, barely above the normal cut-off value.

III. CLINICAL EVALUATION OF IMMUNOASSAY FOR CIRCULATING LCAT

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35 peroxidase-conjugated MAb DF-L1 for 1 h. The plates were
 the plates were washed and incubated with horseradish
 (1:51) were added to the wells. After a 1 h incubation,
 (PBS) for 1 h, washed, and samples containing antigen
 serum albumin (BSA) in 0.1 M phosphate buffered saline
 buffer at pH 8.5). The wells were blocked with 5% bovine
 30 well microtiter culture plates (0.1 M NaClO₃/0.5 M NaCl
 monoclonal antibody DF-L1 (50 μg/ml) was adsorbed to 96
 Tercumo Medical Corporation, Elkhorn, MD. Briefly,
 the method set forth below. The kits are manufactured by
 25 using LCAP ELISA kits as described in detail below, using
 LCAP ELISA. Circulating LCAP levels were assayed
 considered significant.

25 marker, variations in levels of that marker were not
 neither Ig nor Ig was above the cut-off for the respective
 (Tonolini et al., Cancer Res. 48:4107-12, 1988). If
 marker levels were considered significant changes
 20 tumor level. Increases or decreases of >25% in serial tumor
 documented clinical change, and Ig represents the initial
 in which Ig represents the level at the time of first
 15

$$\% \text{ change tumor marker} = \frac{I_f - I_i}{I_i} \times 100$$

calculated as:

10 Percent change in tumor marker (LCAP or CEA) was
 new malignant effusion was considered as progression.
 were considered evaluable, although the appearance of a
 least 60 days. Bone lesions, but not pleural effusions,
 regression nor progression of documented disease for at
 disease. Stable disease was defined as neither
 5 was defined as complete disappearance of known measurable
 50%, lasting for at least 30 days, and complete response
 a decrease in size of a measurable lesion by at least
 dimension of any existing tumor. Response was defined as
 radiographic criteria, or a doubling in the largest
 5

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35 and 40 U/ml, respectively (Table 10). LCAP levels were
 with metastatic lung cancer had LCAP levels >20, 23, 25,
 58 (82%), 55 (77%), 53 (75%), and 30 (42%) of 71 patients
 elevated from non-elevated levels. In the present study,
 we chose 23 U/ml (mean+2 SD) as a cutoff to distinguish
 be 7±8 U/ml, only 1% having LCAP levels above 35 U/ml.
 30 the mean LCAP level in 341 normal subjects was found to
 patients with lung cancer. In the study described above,
 distribution of circulating LCAP levels in

Results

25 NJ, pp. 296-298, 1974.)
 Biosstatistical Analysis, Prentice Hall, Englewood Cliffs,
 differences between proportions (Zar, in J. Zar (ed):
 were performed by calculating the normal deviate (z) for
 (15). Comparisons of two assays in the same population
 20 Wilcoxon test (single factor analysis of variance by ranks)
 different populations was determined using the Kruskal-
 statistics. Comparison of LCAP levels from
 LCAP determinations.

15 Abbott, North Chicago, IL). SCC antigen levels were
 using a microparticulate enzyme immunoassay (IMx SCC,
 determined according to the manufacturer's instructions
 squamous cell carcinoma antigen (SCC antigen) levels were
 10 samples used for LCAP determinations.

Squamous cell carcinoma Antigen Determinations.

CEA levels were determined according
 to the manufacturer's instructions using a microparticulate
 CEA assay. CEA levels were determined according
 (calibrators) provided in the kit.

5 comparison with a curve generated from standards
 stopped with 2 N H₂SO₄; adsorbance was determined at a
 wavelength of 490 nm. LCAP levels were determined by
 developed with o-phenylenediamine and, after 30 min,

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ellevated above 23 U/ml in 28 of 33 (85%) patients with adenocarcinoma, 11 of 13 (85%) patients with squamous cell carcinoma, and 6 of 9 (67%) patients with large cell carcinoma, and 6 of 9 (67%) patients with small cell lung carcinoma (NSCLC), 46 (79%) had LCAP non-small cell lung carcinoma. Of 58 patients with all types of carcinoma of the lung, 9 of 13 (69%) patients also had LCAP values >23 U/ml. Furthermore, 9 of 13 (69%) patients with metastatic small cell carcinoma (SCCLC) or the lung also had LCAP levels >23 U/ml. LCAP levels were >23 U/ml in 15 of 29 (52%) (Table 11). LCAP levels were >23 U/ml in 11 of 29 (38%) and >40 U/ml in 11 of 29 (38%) patients with all histologic subtypes. Elevated levels were observed in 6 of 12 (50%), 3 of 8 (38%), and 1 of 3 (33%) patients with adenocarcinoma, and large cell carcinoma of the lung. Ten of 23 (53%) patients with any type of non-small cell carcinoma had LCAP levels >23 U/ml. In addition, of 5 patients with limited stage small cell carcinoma, 4 (80%) had LCAP levels >23 U/ml and 3 (60%) had LCAP levels >40 had LCAP levels >23 U/ml and 3 (60%) had LCAP levels >40 in these same patients. The reported distribution of CEA in normal subjects (Abbott Package Insert: IMX:CEA, 1990) is compared with that of LCAP in Table 12. We chose a cut-off of 4 ng/ml, which defines roughly 95% of the normal population (including smokers and non-smokers), while 77% of all patients with metastatic lung cancer had LCAP levels >23 U/ml, only 44 of 71 (62%) had elevated CEA levels (Table 12). This difference is statistically significant ($P=0.05$).

ການແພັດທະນາ ໂດຍ ດີເລີ່ມຕົວ

In all patients with metastatic lung cancer, 36 of 71 (51%) had both markers elevated, and only 11 of 71 (15%) had neither. The markers were discordant in 24 of 71 (34%). Sixteen of these 24 (23% of total 71) had only an elevated LCAP level, and 8 of the 24 (11% of total 71) had only an elevated CEA level. In total, 60 (85%) had either LCAP >23 U/ml or CEA >4 ng/ml (Table 12). Combining the two assays was significantly more sensitive than using CEA alone for all patient with metastatic disease (p<0.05). Combining the two assays also increased the sensitivity compared to LCAP alone, but this increase was not statistically significant. Thus, this increase was not statistically significant. Therefore, although LCAP is generally the most sensitive assay for metastatic lung cancer, it is not as effective as two assays combined.

LCAP levels were compared with CEA levels in patients with different sites of disease (Table 13). LCAP levels were more common only elevated than CEA levels in patients without organ involvement (data not shown). Moreover, while the two assays had similar sensitivity in patients with liver metastases, LCAP was significantly more sensitive in patients who did not have liver metastases. Of these 62 patients, 49 (79%) had LCAP levels >23 U/ml, while only 36 (58%) had CEA levels >4 ng/ml (p<0.05) (Table 13). Furthermore, combining the two assays increased sensitivity to 85%, which was significant superior to CEA alone, but not to LCAP alone. Thus, neither assay was very sensitive in patients with minimal metastatic disease (for example,

metastases only to regional lymph nodes), and both assays were equally sensitive in patients with liver metastases. However, LCAP levels were more commonly elevated in patients with liver metastases than in patients with other metastases (Table 14). The overall sensitivity of LCAP was 85%, while CEA sensitivity was 71%.

Thus, LCAP is a useful marker for metastatic lung cancer, especially in patients with liver metastases. It is more sensitive than CEA, and it can be used alone or in combination with CEA to improve diagnostic accuracy.

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patients with distant metastatic disease in whom liver metastases had not yet been detected. In comparison of LCA and SCC antigen levels. In patients with squamous cell carcinoma, LCA levels were compared with SCC antigen. An SCC antigen level of 3 ng/ml was chosen as a cut-off, since previous studies have demonstrated that 95% of a normal population have squamous cell carcinoma of the lung had SCC antigen levels > 3 ng/ml, compared to 77% who had elevated LCA levels (data not shown) ($p < 0.001$). SCC antigen levels ($p < 0.001$). Only 2 of 13 (15%) patients with SCC antigen levels below this level (Body et al., Cancer 65:1552-6, 1990). Only 2 of 13 (15%) patients with squamous cell carcinoma of the lung had SCC antigen levels > 3 ng/ml, compared to 77% who had elevated LCA levels (data not shown) ($p < 0.001$). SCC antigen levels ($p < 0.001$). Correlation of serial LCA levels with clinical course in patients with lung cancer. Changes in LCA levels determined in serial samples were correlated with clinical course (*Table 14*). Overall, serial LCA levels correlated with clinical evaluation of disease course in 21 of 49 (43%) patients. In contrast, serial CEA levels correlated in only 15 of 49 (31%). LCA levels increased correlated in 12 of 25 (48%) patients with progressive disease and decreased in 5 of 7 (71%) patients with disease

In 4 of 17 (23%) patients whose clinical course remained stable over at least 60 days, LCA levels did not change by $\pm 25\%$. Of interest, in the remaining 13 stable patients, LCA levels increased by $> 25\%$ in 4 patients who were found to have progressive disease at the next clinical evaluation. These data indicate that in some patients whose disease was perceived to be clinically stable, increasing LCA levels predicted that

30 subsequent clinical progression.

In 4 of 17 (23%) patients whose clinical course responded to therapy. Combining the two assays improved the correlation with clinical course in 25 patients with progressive disease. In 16 (64%), either serial LCA or CEA levels increased by $> 25\%$ above the respective cutoff.

In 4 of 17 (23%) patients whose clinical course responded to therapy. Combining the two assays improved the correlation with clinical course in 25 patients with progressive disease. In 16 (64%), either serial LCA or CEA levels increased by $> 25\%$ above the respective cutoff.

25 In 4 of 17 (23%) patients whose clinical course remained stable over at least 60 days, LCA levels did not change by $\pm 25\%$. Of interest, in the remaining 13 stable patients, LCA levels increased by $> 25\%$ in 4 patients who were found to have progressive disease at the next clinical evaluation. These data indicate that in some patients whose disease was perceived to be clinically stable, increasing LCA levels predicted that

30 subsequent clinical progression.

SUMMARY STATEMENT

Monitoring of serial LCAP levels post-operatively in patients with primary, non-small cell lung cancer. detectable disease at that time. 25
With newly diagnosed primary NSCLC who underwent complete resection of tumor (Fig. 13). In 7 of these patients, LCAP levels were elevated prior to complete resection of their tumor. Post-operatively, LCAP levels fell on a daily basis. Four patients had pre-operative LCAP levels <50 U/ml, and serial LCAP values fell below the cut-off of 23 U/ml within 2 days. Three other patients had pre-operative LCAP levels between 80 and 100 U/ml, and none of these fell within the normal range within 4 post-operative days. 30

Two other patients in whom LCAP spikes were observed had limited stage small cell carcinoma (Fig. 12B, 12C). Both had pretreatment LCAP levels <30 U/ml, and both achieved a partial response with chemotherapy. In both cases, dramatic rises in LCAP levels were observed during adjuvant radiation therapy to the primary site and the CNS, lasting 4 months or longer. LCAP levels returned to baseline following completion of therapy, and the patients were found to be free of disease at that time.

In 3 patients who responded to therapy, serum LCAP levels exhibited a spike, defined as a dramatic increase followed by a decrease to, or nearly to, baseline (Fig. 12). One of these patients had Stage IIIA adenocarcinoma and was treated with three cycles of combination chemotherapy followed by radiation therapy to the primary site, resulting in a partial response (Fig. 5). During the first cycle of chemotherapy, LCAP levels rose from a baseline of 21 U/ml to a peak of 59 U/ml and then fell to levels slightly above 30 U/ml. Unfortunately, this patient died from post-operative complications, so that long term follow-up was unavailable.

35

by washing the microtiter test wells. Enzyme substrate temperature incubation. Unreacted conjugate is removed then added to the test wells for a one hour room LCAp (DF-L1) conjugated with horseradish peroxidase is by washing the microtiter test wells. Monoclonal anti- room temperature ($18^{\circ}\text{C}-25^{\circ}\text{C}$). Unbound antigen is removed 30 then added to the test wells for a one hour room monoclonal anti-LCAp antibody (DF-L1) for one hour at microtiter test wells that have been coated with In this test for circulating LCAp, serum is incubated in Prinципle of the Test

25

IV. IMMUNOASSAY KIT

and 233 U/ml in 2 patients.

complaints of rheumatoid arthritis, with levels of 210 highest LCAp levels in patients with pulmonary cancer were found in patients with pulmonary 20 11 (45%) patients with chronic bronchitis (range 4-65). pulmonary disease (COPD) (range 9-64 U/ml), and in 5 of in 18 of 33 (55%) patients with chronic obstructive levels were <50 U/ml. Of note, LCAp levels were >23 U/ml 15 levels ranged as high as 233 U/ml, although generally of patients with benign pulmonary disease (Table 15). levels. LCAp levels were also >23 U/ml in 53% (66/125) of 8 patients with gastric cancer, none had elevated LCAp cancer and 3 of 17 (18%) patients with prostate cancer. ovarian cancer, 14 of 26 (54%) patients with pancreatic patients with breast cancer, 26 of 60 (43%) patients with 10 104 (30%) patients with colon cancer, 8 of 14 (57%) tissues (Fig. 14). LCAp levels were >23 U/ml in 31 of with metastatic malignancies of non-lung epithelial circulating LCAp levels were also studied in patients and in patients with benign pulmonary disease. 5

LCAp levels in patient with non-lung malignancies which was calculated during the postoperative decubitus to operative days. The mean ($\pm\text{SD}$) circulating half-life, baseline, was 4.473 days.

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¹⁵ ²⁰ ²⁵ ³⁰

(H_2O_2) and chromophore (ortho-phenylendiamine) are added and allowed to react for thirty minutes at room temperature. The reaction is stopped by the addition of 2N H_2SO_4 . The absorbance of LCAp is directly proportional to the concentration of LCAp. A point-to-point calibrator curve is constructed by plotting the absorbance vs. the dose of LCAp. The spectrophotometer at a wavelength of 492 nm. The absorbance is determined by the unknowns is determined in a suitable technique and the serum should be removed as soon as possible. Handle all samples as if capable of transmitting infectious diseases such as hepatitis B viruses or HIV (AIDS). If the test is to be run within 2 days, the sample should be stored frozen (at least -20°C). For a longer delay between collection and assay, the sample can be stored in a refrigerator (2 - 8°C). Aspecific sample can be stored in a refrigerator (2 - 8°C). Upon receipt, all reagents should be stored at 2 - 8°C. The 20x Wash Concentrate should be stored at room temperature (18-25°C). Do not mix different lots of individual reagents within an assay.

LCAp TEST WELL STRIPS:

12 strips (1 x 8 wells) in frame. Mouse anti-LCAp (monoclonal) immobilized on microtiter strip wells. Keep pouch tightly closed during storage.

The wells were prepared as follows: purified DF-L1 antibody (15 μ g/ml, 0.15 ml/well) in 0.05M aspartic acid and L1 antibodies were prepared as follows: purified DF-L1 antibody (15 μ g/ml, 0.15 ml/well) in 0.05M aspartic acid DF-L1 antibody (15 μ g/ml, 0.15 ml/well) in 0.05M aspartic acid.

Procedure

A. Sample Collection and Storage

B. Reagents Supplied - sufficient for 96 tests

C. Upon receipt, all reagents should be stored at 2 - 8°C. Upon receipt, all reagents should be stored at 2 - 8°C.

D. Reagents Supplied - sufficient for 96 tests

E. Sample Collection and Storage

F. Procedure

G. Sample Collection and Storage

H. Reagents Supplied - sufficient for 96 tests

I. Reagents Supplied - sufficient for 96 tests

J. Reagents Supplied - sufficient for 96 tests

K. Reagents Supplied - sufficient for 96 tests

L. Reagents Supplied - sufficient for 96 tests

M. Reagents Supplied - sufficient for 96 tests

N. Reagents Supplied - sufficient for 96 tests

O. Reagents Supplied - sufficient for 96 tests

P. Reagents Supplied - sufficient for 96 tests

Q. Reagents Supplied - sufficient for 96 tests

R. Reagents Supplied - sufficient for 96 tests

S. Reagents Supplied - sufficient for 96 tests

T. Reagents Supplied - sufficient for 96 tests

U. Reagents Supplied - sufficient for 96 tests

V. Reagents Supplied - sufficient for 96 tests

W. Reagents Supplied - sufficient for 96 tests

X. Reagents Supplied - sufficient for 96 tests

Y. Reagents Supplied - sufficient for 96 tests

Z. Reagents Supplied - sufficient for 96 tests

- 5 CAP CALIBRATORS:
acid coating buffer, pH 3.8, was coated on micro titer strips (Nunc), followed by post-coating with 1% bovine serum albumin (BSA) in 0.01M phosphate buffer, pH 7.4, and drying under nitrogen.
- 10 LCAp CONTROLS:
1x0.8 mL each of four (4) concentrations of LCAp M NaCl; 10% fetal calf serum (GIBCO); and 0.1% sodium buffered solution [0.01M sodium phosphate, pH 4.0; 0.15M NaCl; 2% normal mouse serum; 0.02% merthiolate. serum with 1% sodium azide.
- 15 LCAp SAMPLE DIILUENT: 1 x 25 mL 0.01M phosphate, pH 7.4; 1x0.1mL each of 25 u/mL and 75 u/mL LCAp in human serum with 1% sodium azide].
- 20 ANTI-LCAp HRP CONJUGATE:
1 x 50 mL of 0.2M phosphate, pH 7.4; 0.3M NaCl; 0.2% thimerosal; and 2% tween 20 (Sigma).
- 25 COLORIMETRIC REAGENT:
(DF-L1) in a ratio of 0.8 to 1.2 HRP molecules per molecule of antibody, in a 0.05M Hepes-buffered solution (pH 7.4) with 5% normal mouse serum and 0.02% thimerosal.
- 30 ODD REAGENT:
Hydrogen peroxide.
- 1 x 5 tablets. Each tablet contains 15 mg o-phenylenediamine dihydrochloride and 140 mg excipient.
- 35 STOP REAGENT:
1 x 10 mL 2 N sulfuric acid. Avoid contact with eyes and skin.
- C. Materials Required but Not Supplied

E. Assay Protocol

- 30 3. COLOR DEVELOPING SOLUTION: Prepare sufficient
COLOR DEVELOPING SOLUTION prior to use by dissolving one ODD TABLET in 5 mL of
COLOR DEVELOPING SOLUTION fresh 10 minutes prior
stored at room temperature.
- 25 2. WASH BUFFER (20X): Prepare WASH BUFFER by diluting
50 mL with distilled or deionized water to a final
volume of 1 Liter. Allow any crystals in the
concentrate to dissolve at room temperature
followed by thorough mixing before dispensing a
partial volume. The diluted wash buffer may be
partially used by dissolving crystals in the
assay diluent.
- 20 1. SAMPLE and CONTROL PREPARATION (1:51): Dilute each
sample and control 1:51 with SAMPLE DILUENT by
mixing 10 μ L of sample or control with 0.5 mL of
diluent.
- 15 D. Reagent Preparation
Microtiter: A suitable microtiter plate colorimeter
or spectrophotometer that can measure
absorbance at 492 nm.
Miscellaneous: Microtubes (for handling volumes of
0.5 mL)
- 5 Pipets: 10 μ L - to prepare dilution of samples
and controls 0.5 μ L - to prepare
dispense CONJUGATE, COLOR DEVELOPING
SOLUTION and STOP REAGENT.
Plate Washer: Capable of washing an 8 well strip or 96
well plate.
Vortex Mixer: Vortex mixer or equivalent
50-200 μ L multi-channel pipette - to
dispense samples and controls.
Plate Reader: A suitable microtiter plate colorimeter
or spectrophotometer that can measure
absorbance at 492 nm.
- P. PLATE WASHER: Capable of washing an 8 well strip or 96
well plate.
- Mixer: Vortex mixer or equivalent
SOLUTION and STOP REAGENT.
- P. PLATE WASHER: Capable of washing an 8 well strip or 96
well plate.
- 10 Plate Washer: Capable of washing an 8 well strip or 96
well plate.
- 15 1. SAMPLE and CONTROL PREPARATION (1:51): Dilute each
sample and control 1:51 with SAMPLE DILUENT by
mixing 10 μ L of sample or control with 0.5 mL of
diluent.
- 20 2. WASH BUFFER (20X): Prepare WASH BUFFER by diluting
50 mL with distilled or deionized water to a final
volume of 1 Liter. Allow any crystals in the
concentrate to dissolve at room temperature
followed by dissolving crystals in the
assay diluent.
- 25 3. COLOR DEVELOPING SOLUTION: Prepare sufficient
COLOR DEVELOPING SOLUTION prior to use by dissolving one ODD TABLET in 5 mL of
COLOR DEVELOPING SOLUTION fresh 10 minutes prior
stored at room temperature.

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1. Allow all reagents to reach room temperature prior to use. Mix all reagents thoroughly prior to use.

2. Remove the antibody coated plate from the package of LCAP TEST WELL STRIPS. Return unused LCAP TEST WELL STRIPS to the original pouch provided and seal cut end with tape.

3. Dilute all samples and controls 1:51 in tubes of SAMPLE DILUENT. Vortex briefly to thoroughly (microtubes) by adding 10 μ L of sample to 0.5 mL of SAMPLE DILUENT. Vortex briefly to thoroughly dilute all wells needed for the run with mix.

4. Wash: Fill all wells needed for the run with diluted WASH BUFFER. Allow to soak for 15 ± 5 minutes. Aspirate the plate and wash three times with a suitable microtiter plate washer (1 x 8, or 96 well) or washing manifold (8 or 12 pads to remove any excess wash solution. Invert the plate and tap on absorbent channels). Invert the plate and tap on absorbent as much residual liquid as possible.

5. Note: CALIBRATORS are ready to use: no dilution is necessary.

6. Pipet 100 μ L of the diluted samples in duplicate into the bottom of the appropriate wells.

7. Incubate at control led room temperature (18 to 25°C) for one hour ± 5 minutes. Start timing.

8. After addition of last sample, repeat addition of last sample.

9. Usin a multi-channel pipet, add 200 μ L of ANTI-LCAP-HRP CONJUGATE into the bottom of each well.

10. Incubate at control led room temperature for one hour ± 5 minutes.

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- 30 results are shown in Table 18.
3. Derivative unknowns from the calibrator curve as illustrated in Table 16. It is advisable to run controls of known values.
- NOTE: A correction for specimen dilution is not necessary since the calibrators are provided in ready-to-use form.
4. Samples with LCAP concentrations greater than 200 UNITS/mL CALIBRATOR should be further diluted into the assay range with SAMPLE DILUENT. The obtained value should be multiplied by this dilution factor to obtain the unit value.
5. Limitations of the Procedure
1. Samples with LCAP concentrations greater than the 200 UNITS/mL CALIBRATOR should be further diluted into the assay range with SAMPLE DILUENT. The although the assay performance has been established between 18° and 25°C, the absorbance at the 200 unit calibrator has been set at 22°C.
- 15 ambient temperatures above or below 22°C may affect the absorbance.
2. Although the assay performance has been established between 18° and 25°C, the absorbance at the 200 unit calibrator has been set at 22°C. of the 200 unit calibrator has been set at 22°C.
- 20 three unknown samples were diluted with SAMPLE DILUENT and assayed for LCAP concentration. The 1:1 sample was used to define the expected values on dilution. (Note: Each sample was previously diluted known quantities of LCAP were added to four diluted specimens. Recovery was calculated by subtracting the endogenous level from the assayed value, dividing by quantity added, and multiplying by 100).
- 25 b. Recovery
- 1:51). The results are shown in Table 17.
2. Dilution Linearity
- Specifc Performance Characteristics
- 200 UNITS/mL CALIBRATOR should be further diluted into the assay range with SAMPLE DILUENT. Three unknown samples were diluted with SAMPLE DILUENT and assayed for LCAP concentration. The 1:1 diluent and assayed for LCAP concentration. The 1:1 dilution was used to define the expected values on dilution. (Note: Each sample was previously diluted known quantities of LCAP were added to four diluted specimens. Recovery was calculated by subtracting the endogenous level from the assayed value, dividing by quantity added, and multiplying by 100).
- 30 results are shown in Table 18.

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- 35 prepared by immunization of mice or other animals with
producing LCA-P-specific monoclonal antibodies may be
claims set forth below. For example, hybridomas
other embodiments of the invention are within the
Other Embodiments
- 30 consistently less than or equal to 1.60 units/ml.
the same calibrator curve. The LCA-P value obtained was
unit CALIBRATOR, and interpolating the LCA-P value from
two standard deviations to the mean absorbance for the 0
mean and standard deviations for the calibrators, adding
running the calibrator curve six times, calculating the
The sensitivity of the LCA-P assay was determined by
E. Sensitivity
- 25 quantity added and multiplying by 100.
the apparent LCA-P value (corrected for endogenous) by the
reactivity shown in Table 22 was calculated by dividing
assayed to obtain the endogenous LCA-P value. The cross
concentration indicated in Table 22. Unspiked serum was
assay was determined by adding the marker up to the
2. The effect of other tumor markers on the LCA-P
to the indicated levels of these substances.
15 detectable interference was observed in samples having up
bilirubin or rheumatoid factor. As shown in Table 21, no
serum samples containing, respectively, triglycerides,
1. Varying levels of LCA-P were added to human
D. Specificity
20 statistics were obtained by assaying the five serum
samples in replicates of four in each of three assays
over three days (Table 20).
25 the precision of the assay was determined by assaying each
of five human serum samples in replicates of four (Table
19).
30 1. Inter-assay variation. The "between" run
2. Inter-assay variation. Data for determining
C. Precision
35

technology available to those of ordinary skill in the art of making such hybrids (see, e.g., Murphy, U.S. Pat. No. 4,675,382, and Chaudhary et al., Proc. Natl. Acad. Sci. USA 84:4538-4542, 1987; each of which is herein incorporated by reference). The DNA sequence encoding the LCAP-binding portion of the immunotoxin would be based upon the variable light-chain (V_L) amino acid sequence and the variable heavy-chain (V_H) sequence of an LCAP-specific antibody of the invention; using the method of Bird et al., Science 242:423-426, 1988, a DNA sequence of V_L joined to the V_H by a linker peptide would be constructed and linked to a DNA sequence encoding the protein toxin (or a toxic portion thereof, as taught by, for example, Murphy U.S. Pat. No. 4,675,382). Such manipulations would be routine to one of ordinary skill in the art of genetic engineering, given the disclosure set forth herein. The resulting immunotoxin could be formulated for use as an anti-cancer agent, following procedures standard to the field of pharmacology.

An LCAP-specific monoclonal antibody can produce an imaging agent useful for detecting and localizing LCAP-expressing tumors *in vivo*. Methods of attaching such labels to antibodies are well known in the art, and can be readily accomplished without undue experiment. The potential usefulness of such specific tumor cells into an immunocompetent host (such as a nude mouse) and determine whether or not the tumor produced by such implanted cells.

The vaccine of the invention includes the LCAP protein core, or an antigenic fragment of the LCAP has been removed, or an anti-geneic fragment of the LCAP protein core from which some or all of the carbohydrate protein chain has been removed, or an anti-geneic fragment of the LCAP protein core which retains some or all of the carbohydrate chain.

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25 fragments.

manipulated to encode and express defined peptide
commun. 165:644-649, 1989, and then genetically
6971, 1989; and Abe and Kufe, Biochem. Biophys. Res.
85:2320-2323, 1988; Merlo et al., Cancer Res. 49:6966-
described in Saidiqui et al., Proc. Natl. Acad. Sci. USA
cloned and sequenced using methods similar to those
example, DNA encoding the LCAP core protein could be
by genetic engineering, using standard methods. For
protein could be produced enzymatically, chemically, or
immune system. Peptide fragments of the LCAP core
expressing tumor cells for attack by the patient's own
antibodies so raised would target any such LCAP-
carbohydrate makew up than LCAP of normal cells), the
express an LCAP antigen with a slightly different
their altered metabolism relative to normal cells may
relativeley exposed on certain tumor cells (which due to
immune response. To the extent that these sites are also
can, when injected as a vaccine in a human, induce an
naturally-occurring glycoprotein, and which therefore
uncovers antigenic sites which are hidden in the
all of the carbohydrate from the LCAP protein core
not inherently immunogenic in humans, removing some or
the circulation of most normal individuals and thus is
human glyccoprotein found (at relatively low levels) in

Table 1
Reactivity of Mabs DF-L1 and DF-L2 with human tumor cell lines

Tumor cell line	Reactivity*	
	MAb DF-L1	MAbDF-L2
Lung		
CALU-3	+++	+++
SK-MES	++	+++
A-549	+++	+++
CALU-1	++	+++
Breast		
MCF-7	+++	++
ZR-75-1	+++	+++
BT-20	+++	+++
Ovarian		
OVCAR	++	-
OV-S	-	-
OV-D	++	+
Melanoma		
A-374	-	-
Leukemia		
U-937	-	-
HL-60	-	-

*++, strong; ++, moderate; +, weak; -, no reactivity.

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Table 2
Immunoperoxidase staining of lung tumor tissues with MAb DF-L1

Histology	Tumors positive/tested	% cells positive	Pattern*	
			A	C
Adenocarcinoma	6/6	40-100	+++	++
Epidermoid	4/4	20-75	+++	++
Small cell	0/5	0	-	-

*A, apical; C, cytoplasmic; +++, intense staining; ++, moderate staining; +, weak staining; -, no staining

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*A, apical; C, cytoplasmic; ++, intense staining; +, moderate staining; -, no staining.

Tissues	Specimens	Cell type	positive/tested	positive	A	C
Normal						
Spleen	0/1	0/2	0/2	0/1	Ovary	Testis
Heart	0/2	0/1	0/2	0/2	Cervix	
Liver	0/3	0/1	0/2	0/2		
Muscle	0/2	0/2	0/5	0/5	Lymphoid	Bowel
Cartilage	0/2	0/2	0/5	0/5		
Bone	0/1	0/1	0/1	0/1		
Endothelium	0/5	0/5	2/2	2/2	Breast	Kidney
Thyroid	2/2	Follicular	4/4	4/4	Breast	Kidney
Pancreas	5/5	Ducts	3/3	3/3	Stomach	
Lung	5/5	Pneumatoctyes	1/2	1/2		
Tumor	10/10	Mucus-producing	5/5	5/5	Ovary	Melanoma
					Sarcoma	Lymphoma

Table 3 Immunooperoxidase staining of normal tissues and non-lung tumors with MAb DF-L1

Table 5					for the detection of circulating LCAp Comparison of assay formats			
Mean LCAp levels with tracer MAb					DF-L1-HRP			
Capture MAb subjects patients subjects patients					Normal Cancer Normal Cancer			
DF-L1	7	92	8	65	DF-L2	1	7	51
DF-L4	NE**	NE	8	67	DF-L4	NE**	NE	6
*	Means units/ml of 5 samples.							** NE: not evaluated.

Table 4					Analyses of MAb DF-L1 and DF-L2 binding sites			
Dot blot*					ELISA (% of control)			
Treatment					DF-L2	DF-L1	DF-L2	
None	+	+	100	100	-	-	-	34
Trypsin	+	+	77	77	-	-	100	16
Protease	-	-	3	3	+	+	95	91
Neuraminidase	-	-	2	2	+	+	95	91
Periodate	+	+	56	56	+	+	100	100
NAOA/borohydride	+	+	79	79	-	-	-	79

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0 units/ml	0.061 ± 0.006	9.52	200 units/ml	2.442 ± 0.105	4.29	Specimen 3	27.7	± 2.11	7.61	
20 units/ml	0.343 ± 0.031	8.90	100 units/ml	1.374 ± 0.072	5.25	Specimen 2	56.2	± 2.32	4.13	
100 units/ml	0.343 ± 0.031	8.90	200 units/ml	2.442 ± 0.105	4.29	Specimen 1	87.8	± 3.73	4.25	
200 units/ml	2.442 ± 0.105	4.29	Specimen 1	87.8	± 3.73	4.25	Specimen 1	87.8	± 3.73	4.25

Table 7 Intrassay reproducibility of LCAP assay
 Sample Means ± SD at A₄₉₀ Coefficient of variation (%) (n = 7)

0 units/ml	0.05 ± 0.002	3.90	200 units/ml	2.64 ± 0.070	2.67	Specimen 1	1.34 ± 0.060	4.14	85.6
20 units/ml	0.38 ± 0.020	5.04	100 units/ml	1.55 ± 0.090	5.57	Specimen 2	0.85 ± 0.050	5.74	52.1
100 units/ml	0.38 ± 0.020	5.04	200 units/ml	2.64 ± 0.070	2.67	Specimen 3	0.46 ± 0.020	4.88	25.5
200 units/ml	2.64 ± 0.070	2.67	Specimen 1	1.34 ± 0.060	4.14	Specimen 2	0.85 ± 0.050	5.74	52.1

Table 6 Intrassay reproducibility of LCAP assay
 Sample Mean ± SD at A₄₉₀ Coefficient of variation (%) (units/ml) (n = 12)

0 units/ml	0.05 ± 0.002	3.90	200 units/ml	2.64 ± 0.070	2.67	Specimen 1	1.34 ± 0.060	4.14	85.6
20 units/ml	0.38 ± 0.020	5.04	100 units/ml	1.55 ± 0.090	5.57	Specimen 2	0.85 ± 0.050	5.74	52.1
100 units/ml	0.38 ± 0.020	5.04	200 units/ml	2.64 ± 0.070	2.67	Specimen 3	0.46 ± 0.020	4.88	25.5
200 units/ml	2.64 ± 0.070	2.67	Specimen 1	1.34 ± 0.060	4.14	Specimen 2	0.85 ± 0.050	5.74	52.1

Table 8

Recovery of LCAP after addition to plasma

Specimen	Endogenous LCAP (units/ml)	LCAP added (units/ml)	Expected (units/ml)	Observed (units/ml)	Recovery (%)
1	1.6	179.0	180.6	174.1	96.4
2	4.6	141.0	145.6	144.9	99.5
3	7.9	94.0	101.9	100.0	98.0
4	12.6	45.0	57.6	61.0	106.0

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		Bilirubin (mg/dl)		Rheumatoid factor		Triglycerides (mg/dl)	
Substance and LCA added	Recovery (%)	Concentration (units/ml)		Interference studies			
16.3	109.2	128.3	128.3	110.6	19.5	3.0	99.4
19.5	110.6	128.3	128.3	106.0	128.0	120.0	104.2
19.5	110.6	128.3	128.3	120.0	120.0	120.0	87.2
22.1	108.2	128.3	128.3	120.0	120.0	120.0	92.9
22.1	108.2	128.3	128.3	120.0	120.0	120.0	92.9
96.9	96.9	120.0	120.0	120.0	120.0	120.0	98.0
96.9	96.9	120.0	120.0	120.0	120.0	120.0	104.2
536	103.1	128.3	128.3	128.0	128.0	128.0	103.4
536	103.1	128.3	128.3	128.0	128.0	128.0	105.9
415	106.0	128.3	128.3	128.0	128.0	128.0	106.0
415	106.0	128.3	128.3	128.0	128.0	128.0	105.0
426	105.0	128.3	128.3	128.0	128.0	128.0	105.0
345	112.0	128.3	128.3	128.0	128.0	128.0	112.0
345	112.0	128.3	128.3	128.0	128.0	128.0	112.0

Table 10

Distribution of Circulating LCAP Levels in Patients with Metastatic Lung Cancer

No. (%) of Patients with LCAP levels:

Histologic Category	No. Pts.	>15 (U/mL)	>20 (U/mL)	>23 (U/mL)	>25 (U/mL)	>30 (U/mL)	>35 (U/mL)	>40 (U/mL)	Mean LCAP (+SD)
Normal	341	47 (14)	25 (7)	18 (5.3)	15 (4)	8 (2)	4 (1)	1 (<1)	7.2±8
All Lung Cancers	71	63 (89)	58 (82)	55 (77)	53 (75)	46 (65)	38 (54)	30 (42)	114.4±249
All NSCLC	58	54 (93)	49 (84)	46 (79)	45 (78)	39 (67)	35 (60)	28 (48)	133.3±272
Adeno	33	31 (94)	29 (88)	28 (85)	27 (82)	23 (70)	22 (66)	19 (58)	172.5±321
Squamous	13	13 (100)	11 (85)	11 (85)	10 (77)	8 (62)	6 (46)	3 (10)	105.8±244
Large Cell	9	7 (78)	7 (78)	6 (67)	6 (67)	6 (67)	6 (67)	5 (56)	68.4±85
Misc.									
NSCLC	4	4(100)	3 (75)	3 (75)	3 (75)	2 (50)	2 (50)	35.5±33	
SCLC	13	9 (69)	9 (69)	9 (69)	8 (62)	7 (54)	3 (23)	2 (15)	22.8±11.8

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Table 11
Distribution of circulating LCAP levels in Patients with Primary Lung Cancer

No. (%) of Patients with LCAP levels:

Histologic Category	No. Pts.	>15 (U/ml)	>20 (U/ml)	>23 (U/ml)	>25 (U/ml)	>30 (U/ml)	>35 (U/ml)	>40 (U/ml)	Mean LCAP (+SD)
All Lung Cancers	29	24 (83)	22 (76)	15 (52)	14 (48)	12 (41)	12 (41)	11 (38)	39.3±33
All NSCLC	24	19 (83)	17 (74)	10 (43)	10 (43)	9 (39)	9 (39)	9 (39)	37.3±33
Adeno	12	11 (92)	9 (75)	6 (50)	6 (50)	5 (42)	5 (42)	4 (33)	32.8±17
Squamous	8	5 (63)	5 (63)	3 (38)	3 (38)	3 (38)	3 (38)	3 (38)	41.6±50
Large cell	3	3 (100)	3 (100)	1 (33)	1 (33)	1 (33)	1 (33)	1 (33)	43.3±37
SCLC	5	4 (80)	4 (80)	4 (80)	4 (80)	3 (60)	3 (60)	3 (60)	51.6±34

Table 12
Comparison of LCAP and CEA Levels in Patients
with Metastatic Lung Cancer

	No. (%) of Patients with:		
	Either		
	LCAP>23 U/ml or	CEA>4 ng/ml	CEA >4 ng/ml
Normal	(5)	(5)	(5)
All Lung Cancers	71 55 (77)*	44 (62) 38 (66)	60 (85)* 51 (88)*
All NSCLC	58 46 (79)	38 (66)	51 (88)*
Adeno	33 28 (85)	25 (76)	29 (88)*
Squamous	13 11 (85)	7 (54)	12 (92)
Large	9 6 (67)	4 (44)	7 (78)
Misc.			
NSCLC	4 3 (75)	2 (50)	4 (100)
SCLC	13 9 (69)	6 (46)	9 (69)

p<0.05 compared to CEA alone.

Table 13
Comparison of LCAP and CEA Levels
in Patients with Metastatic Lung Cancer by site of Disease

<u>ANY LUNG BONE OR LIVER METASTASES</u>										<u>NO LIVER METASTASES</u>				
No. (%) of Patients with:					No. (%) of Patients with:					Either				
Either		LCAP			CEA		LCAP>23 U/ml		LCAP		CEA		LCAP>23 U/ml	
Histologic Category	No.	Levels >23 Pts. (U/ml)	Levels >4 or (ng/ml)	CEA >4 ng/ml	No.	Levels >23 Pts. (U/ml)	Levels >4 or (ng/ml)	CEA >4 ng/ml	No.	Levels >23 Pts. (U/ml)	Levels >4 or (ng/ml)	CEA >4 ng/ml	No.	Levels >23 Pts. (U/ml)
All	56	44 (79)**	35 (63)	48 (86)*	62	49 (79)	36 (58)	53 (85)*						
All NSC	46	38 (83)**	31 (67)	42 (91)*	52	43 (83)	33 (63)	47 (90)*						
Adeno	28	25 (89)	21 (75)	26 (93)**	29	25 (86)	22 (86)	27 (93)*						
Squam	10	8 (80)	6 (60)	9 (90)	12	11 (92)	6 (50)	11 (92)						
Large Cell	6	4 (67)	3 (50)	5 (83)	8	6 (75)	3 (38)	6 (75)						
Misc.	3	2 (67)	1 (33)	3 (100)	4	3 (75)	2 (50)	4 (100)						
NSC	10	6 (60)	4 (40)	6 (60)	10	6 (60)	3 (30)	6 (60)						
SCC														

*p<0.05 compared to CEA alone.

**0.08<p<0.05 compared to CEA alone.

Table 14

Correlation of Serial LCAP and CEA Levels with Clinical course of Disease

Number (%) of Patients with change in Antigen
Level that Correlates with clinical course^{1,2}

<u>Disease Course</u>	<u>No. of Pts.</u>	<u>LCAP</u>	<u>CEA</u>	<u>LCAP or CEA</u>	<u>Either</u>
Progression	25	12 (48)	8 (32)	16 (64)	
Response	7	5 (71)	4 (57)	5 (71)	
Stable	17	4 (23)	3 (16)	1 (6)	
All	49	21 (43)	15 (31)	22 (45)	

¹For patients with progressive disease, antigen levels increased by >25%. For patients with responding disease, antigen levels decreased by >25%. For patients with stable disease, antigen levels did not increase or decrease by ±25%.

²If antigen level never above cutoff (LCAP>23 U/ml, CEA>4 ng/ml), antigen not considered to correlate regardless of % change.

Table 15

LCAP Levels in Patients with Benign Pulmonary Conditions

<u>Condition</u>	<u>Pts.</u>	<u>>23 U/ml (%)</u>	<u>(Mean \pm SD)</u>	<u>Range</u>
All	125	66 (53)	37.0 \pm 40	3.0 - 233.0
Asthma	6	1 (17)	22.3 \pm 18	5.0 - 58.0
Bronchiectasis	3	2 (67)	55.0 \pm 34	16.0 - 81.0
Bronchitis	13	7 (54)	29.9 \pm 17	4.0 - 65.0
COPD	33	18 (55)	27.2 \pm 17	9.0 - 64.0
Pneumonia	28	16 (57)	31.6 \pm 23	4.9 - 100.0
Pneumothorax	3	1 (33)	54.7 \pm 89	3.0 - 158.0
Pulm. Embolus	4	4 (100)	42.5 \pm 7	35.0 - 52.0
Tuberculosis	17	8 (47)	31.0 \pm 20	4.0 - 78.0
Cystic Fibrosis	4	2 (50)	26.3 \pm 15	13.0 - 49.0
Sarcoidosis	3	2 (67)	88.3 \pm 89	20.0 - 189.0
Misc. 1	11	5 (45)	93.9 \pm 102	3.0 - 233.0

¹Includes Acute Respiratory Insufficiency, Alveolitis, Atelectasis, Bronchial Polyp, Empyema, Hemoptysis, Rheumatoid Lung, silicosis, Pulmonary Edema, Subcutaneous Emphysema.

	Code	Absorb.	Mean	LCAp Absorb.	Units/ml
0 UNIT CAT	S1	0.000	0.000	-	-
20 UNIT CAT	S2	0.288	0.290	0.291	0.291
100 UNIT CAT	S3	1.205	1.217	1.229	1.229
200 UNIT CAT	S4	2.325	2.344	2.362	2.362
Control Level 1	C1	0.287	0.293	0.299	0.299
Control Level 2	C2	1.492	1.505	1.518	1.518
Unknown #1	U1	0.396	0.400	0.404	0.404
Unknown #2	U2	1.350	1.300	1.250	1.250

Typical Data obtained with LCAp Immunoassay Kit

Table 16

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Dilution	Expected Sample (U/ml)	Measured (U/ml)	Correlation (R value)
1:1	1	50.8	0.999
1:2	2	85.8	1.000
1:4	4	42.9	42.8
1:8	8	21.5	20.8
1:16	16	10.7	9.3
1:32	32	155.0	0.997
1:64	64	77.5	85.0
1:128	128	38.8	35.0
1:256	256	19.4	15.0

Table 17

Table 18

Specimen	Endogenous LCAP (U/mL)	Amount Added (U/mL)	Expected (U/mL)	Observed (U/mL)	% Recovery
1	1.6	179.0	180.6	174.1	96.4
2	4.6	141.0	145.6	144.9	99.5
3	7.9	94.0	101.9	100.0	98.0
4	12.6	45.0	57.6	61.0	106.0

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Sample #	n	Mean Units	S.D.	%C.V.
Table 20				
1	3	10.5	1.33	12.64
2	3	28.3	0.53	1.87
3	3	30.9	1.65	5.34
4	3	48.9	2.05	4.20
5	3	94.8	6.38	6.72

Sample #	n	Mean Units	Mean Absorbance	S.D.	%C.V.
Table 19					
1	4	10.2	0.152	0.006	3.99
2	4	28.7	0.372	0.011	2.84
3	4	32.6	0.415	0.013	3.08
4	4	49.4	0.601	0.017	2.81
5	4	93.7	1.158	0.028	2.45

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Substance	Concentration (by Concentration)	Interference	Table 21
INTERFERENCE			
Triglycerides	924.0 mg/dL	Bilirubin None	
		Rheumatoid Factor None	
		Marker [Normal] (LCAp) (U/mL) Assayed [LCAp] (U/mL) % Cross Reaction	
CEA	3 ng/mL	5 ng/mL 13.4	12.2
NSE	22 ng/mL	5 ng/mL 10.1	10.9
CA 19-9	40 U/mL	1000 U/mL 13.0	13.8
CA 125	35 U/mL	500 U/mL 13.0	16.4
CA 15-3	22 U/mL	600 U/mL 14.9	148.4
			22.2
			0.7

SUSAN H. LEE (HARRIS)

1. An essential ally purified preparation of human Lung Cancer-associated Protein (LCAP).
2. The preparation of claim 1, wherein said LCAP is extracted from membranes of human cells.
3. The preparation of claim 1, wherein said LCAP is isolated from a bodily fluid of a person.
4. The preparation of claim 3, wherein said bodily fluid is blood.
5. The preparation of claim 1, wherein said LCAP is secreted by a cell cultured in vitro.
6. A method of making the preparation of claim 1, said method comprising providing a population of cells capable of expressing said LCAP; culturing said population of cells in a medium under conditions which permit said population of cells to express said LCAP; and isolating said LCAP from the membranes of said cells.
7. The method of claim 6, wherein said population of cells is descended from a CALU-3 cell [American Type Culture Collection (ATCC) accession No. HTB 55].
8. Isolating said LCAP from the membranes of said cells to express said LCAP; and
9. Population of cells or from said medium.
10. The method of claim 6, wherein said population of cells is descended from a CALU-3 cell [American Type Culture Collection (ATCC) accession No. HTB 55].

Claims

- 1 8. The method of claim 6, wherein said isolating step comprises the step of contacting said membrane, or said medium with an extract of said membranes, affixed to a matrix material.
- 1 9. The method of claim 8, wherein said antibody is the monoclonal antibody produced by the hybridoma DF-11.
- 1 10. The method of claim 7, wherein said medium comprises galactosamine.
- 1 11. A hybridoma cell which produces an antibody specific for LCAP.
- 1 12. The hybridoma cell of claim 11, wherein said hybridoma is DF-11.
- 1 13. The hybridoma cell of claim 11, wherein said antibody binds to the same determinant on LCAP to which the antibody produced by the hybridoma DF-11 binds.
- 1 14. A monoclonal antibody specific for LCAP.
- 1 15. The monoclonal antibody of claim 14, wherein said monoclonal antibody is produced by the hybridoma DF-11.
- 1 16. The monoclonal antibody of claim 14, wherein said antibody binds to a determinant on LCAP to which the antibody produced by the hybridoma DF-11 binds.
- 2 17. A method of producing a monoclonal antibody specific for LCAP, said method comprising culturing the hybridoma cell of claim 11 in a medium, and isolating said antibody from said medium.

18. A method of detecting LCAP in a biological sample, said method comprising contacting said biological sample with an aliquot containing the monoclonal antibody of claim 14; and detecting the monoclonal antibody and a constituent of said biological sample bearing immunogenicity between said antibody and a monoclonal antibody produced by the hybridoma DF-L1, wherein said biological sample is human serum.
19. The method of claim 18, wherein said complex formation is detected by ELISA.
20. The method of claim 18, wherein said biological sample is human serum.
21. The method of claim 18, wherein said monoclonal antibody is produced by the hybridoma DF-L1.
22. The method of claim 18, wherein said monoclonal antibody binds to a determinant on LCAP to which the monoclonal antibody produced by the hybridoma DF-L1 binds.
23. The method of claim 18, comprising the additional steps of providing a control sample containing a standard amount of LCAP; providing a control sample containing a standard amount of LCAP; and contacting said control sample with a second aliquot containing said control sample with a second amount of LCAP;
24. An immunoassay kit comprising:
- 1 a first reagent comprising a first monoclonal antibody specific for LCAP;
 - 2 a first reagent comprising a first monoclonal antibody specific for LCAP;
 - 3 a second reagent comprising a second monoclonal antibody specific for LCAP;

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- 4 A second reagent comprising an enzyme conjugated to a second monoclonal antibody specific for LCA-P;
- 5 to a second monoclonal antibody specific for LCA-P;
- 6 a third reagent comprising a substrate for said enzyme; and
- 7 the second monoclonal antibodies are specific for the same determinant on LCA-P.
- 8 instructions for using said kit.
- 1 25. The kit of claim 24, wherein said first and second monoclonal antibodies are identical.
- 1 26. The kit of claim 25, wherein said first and second monoclonal antibodies are specific for the same determinant on LCA-P.
- 2 27. The kit of claim 24, wherein one of said first and second monoclonal antibodies binds to the same determinant on LCA-P to which the monoclonal antibody produced by the hybridoma DF-L1 binds.
- 1 28. The kit of claim 27, wherein both of said first and second monoclonal antibodies bind to said determinant.
- 2 29. The kit of claim 24, wherein said enzyme is horseradish peroxidase and said substrate is hydrogen peroxide.
- 1 30. The kit of claim 24, wherein said kit further comprises a fourth reagent comprising a further reagent is a calibrator or control sample.
- 2 31. The kit of claim 30, wherein said fourth reagent is a mixture comprising an LCA-P-specific monoclonal antibody, or an LCA-P-binding fragment thereof, conjugated to a toxin molecule.
- 1 32. An immunoassay comprising an LCA-P-specific monoclonal antibody conjugated to a toxin molecule.

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SERIALIZED FILED

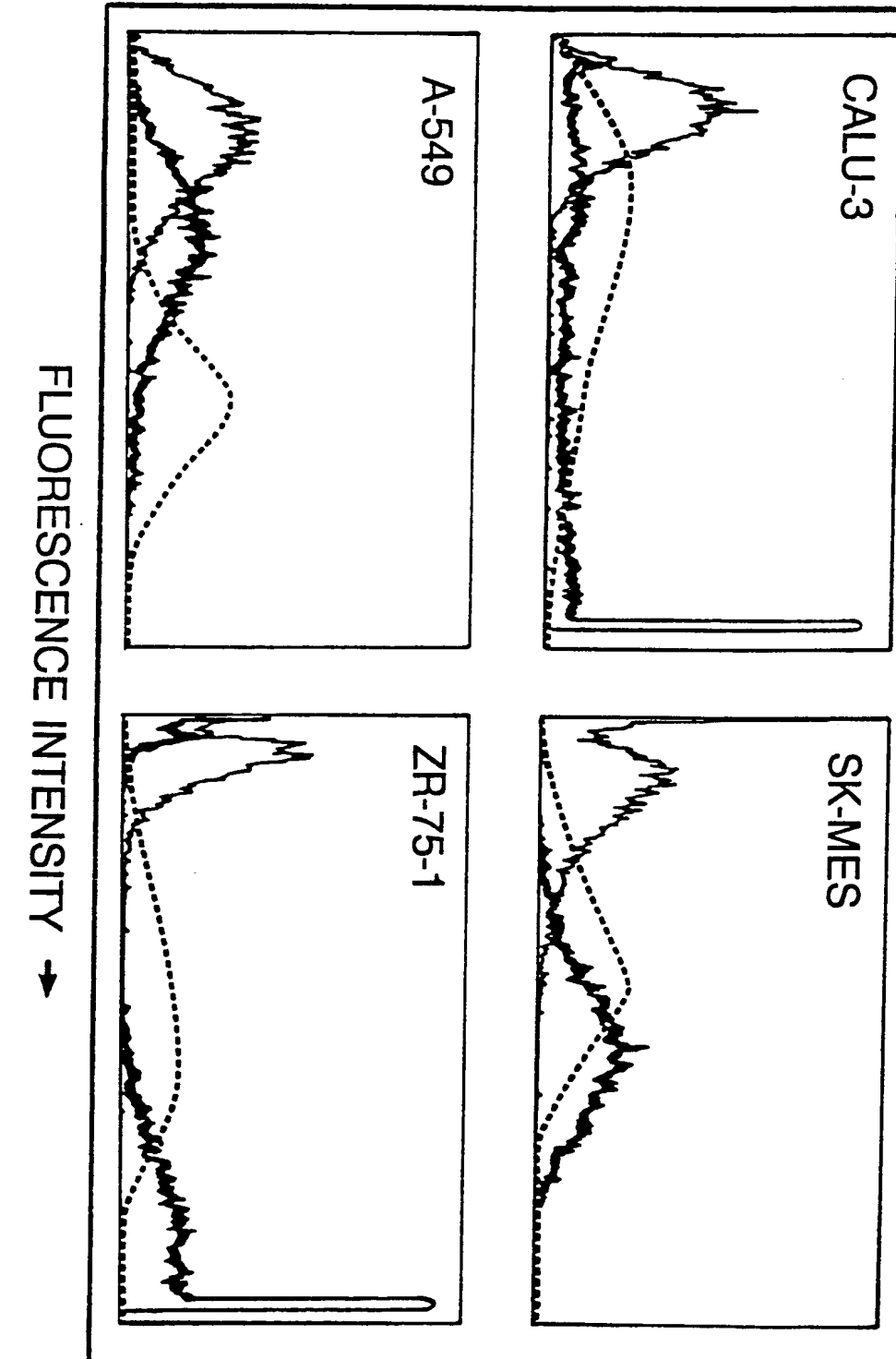
- 1 33. The immunotoxin of claim 32, wherein said toxin molecule is a protein.
- 2 34. The immunotoxin of claim 32, wherein said toxin is chemically conjugated to said monoclonal antibody or said LCAP-bindin fragment.
- 3 35. The immunotoxin of claim 33, wherein said toxin is linked by a peptide bond to said LCAP-bindin fragment, and said immunotoxin is produced by expression of a genetically engineered hybrid DNA molecule.
- 4 36. An imaging agent comprising an LCAP-specific monoclonal antibody, or an LCAP-bindin fragment thereof, linked to a detectable label.
- 1 37. The imaging agent of claim 36, wherein said label is a radionuclide.
- 2 38. The imaging agent of claim 36, wherein said antibody binds to the determinant on LCAp to which the antibody produced by the hybridoma DF-L1 binds.
- 3 39. The imaging agent of claim 36, wherein said antibody is the antibody produced by the hybridoma DF-L1.
- 1 40. A method comprising 11.
- 2 identifying an animal suspected of having a tumor;
- 3 introducing into said animal the imaging agent of
- 4 detecting the presence of said detectable label
- 5 claim 36; and
- 6 bound to a tissue of said animal.
- 7

- 1 41. The method of claim 40, wherein said animal is suspended of having a lung tumor, and said tissue is - 64 -
- 2 42. The method of claim 40, wherein said label is a radioactive and said detection step is accomplished by radiomaging.
- 3 43. A vaccine comprising the LCA P core protein, or a peptide fragment thereof, in a pharmaceutically acceptable carrier.
- 1 44. The vaccine of claim 43, wherein said vaccine additionally comprises an adjuvant.
- 2 45. A method of immunizing a human, said method comprising introducing into said human the vaccine of claim 43.

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CELL
NUMBER

FLUORESCENCE INTENSITY →



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FIG. 2b

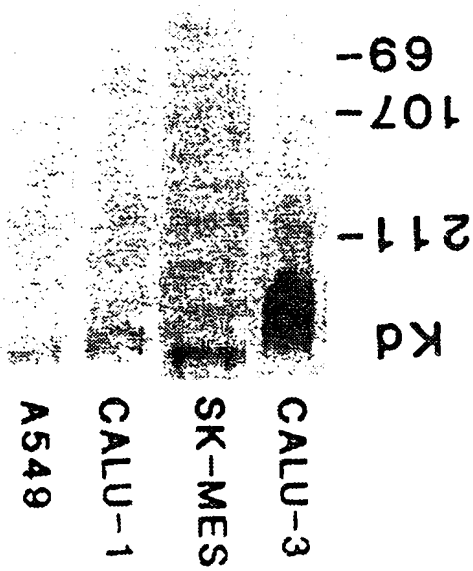


69-
107-
211-

Kd

MCF-7
BT-20
ZR-75-1
OV-A0
OV-S
OVCAR
U-837
HL-60

FIG. 2a



69-
107-
211-

Kd

CALU-3
SK-MES
CALU-1
A549

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FIG. 3b



FIG. 3a



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SURETY BOND
EXPIRED

FIG. 3d



FIG. 3c



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FIG. 5

69-

107-

211-

Kd

Control
Tunicamycin

FIG. 4

69-

107-

211-

Kd

MAb DF-L1
MAb DF-L2
Control

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FIG. 6

-69-

-107-

-211-

Kd

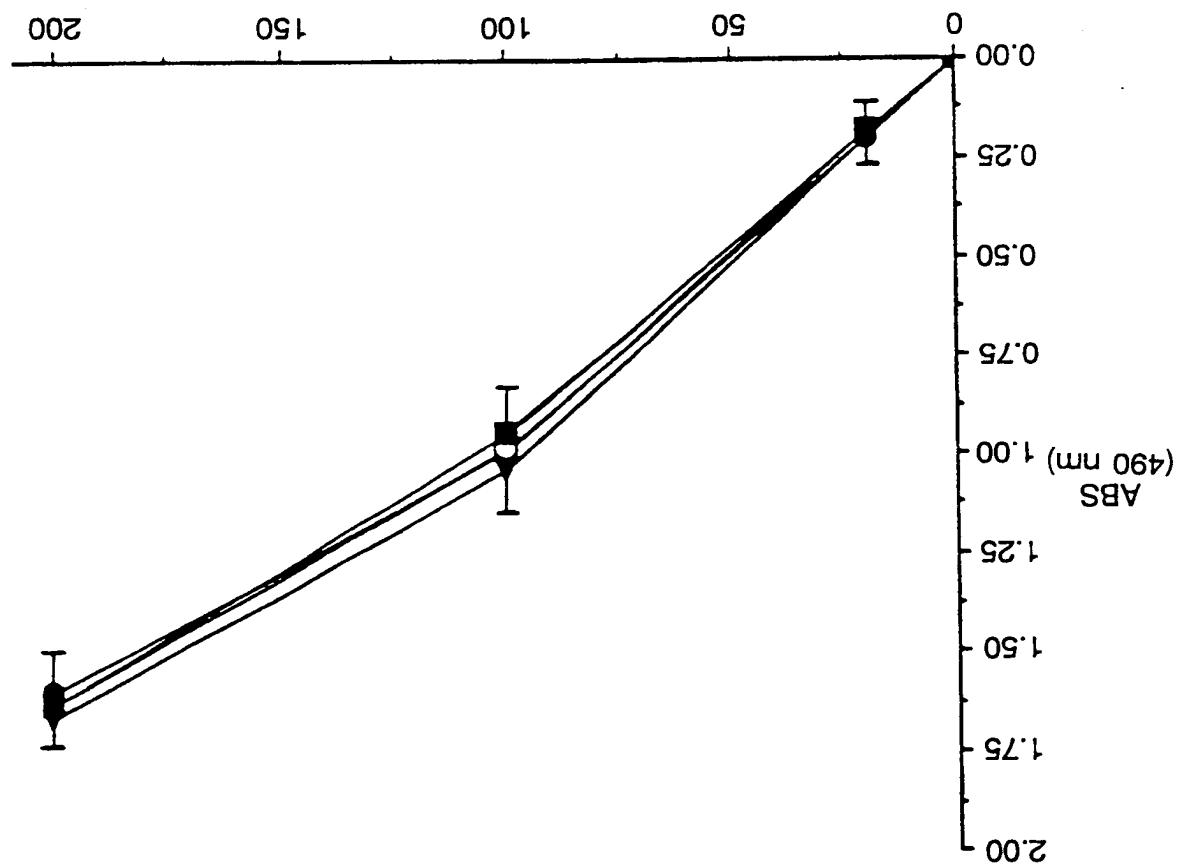
1 2 3 4 5 6 7 8 9

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FIG. 7

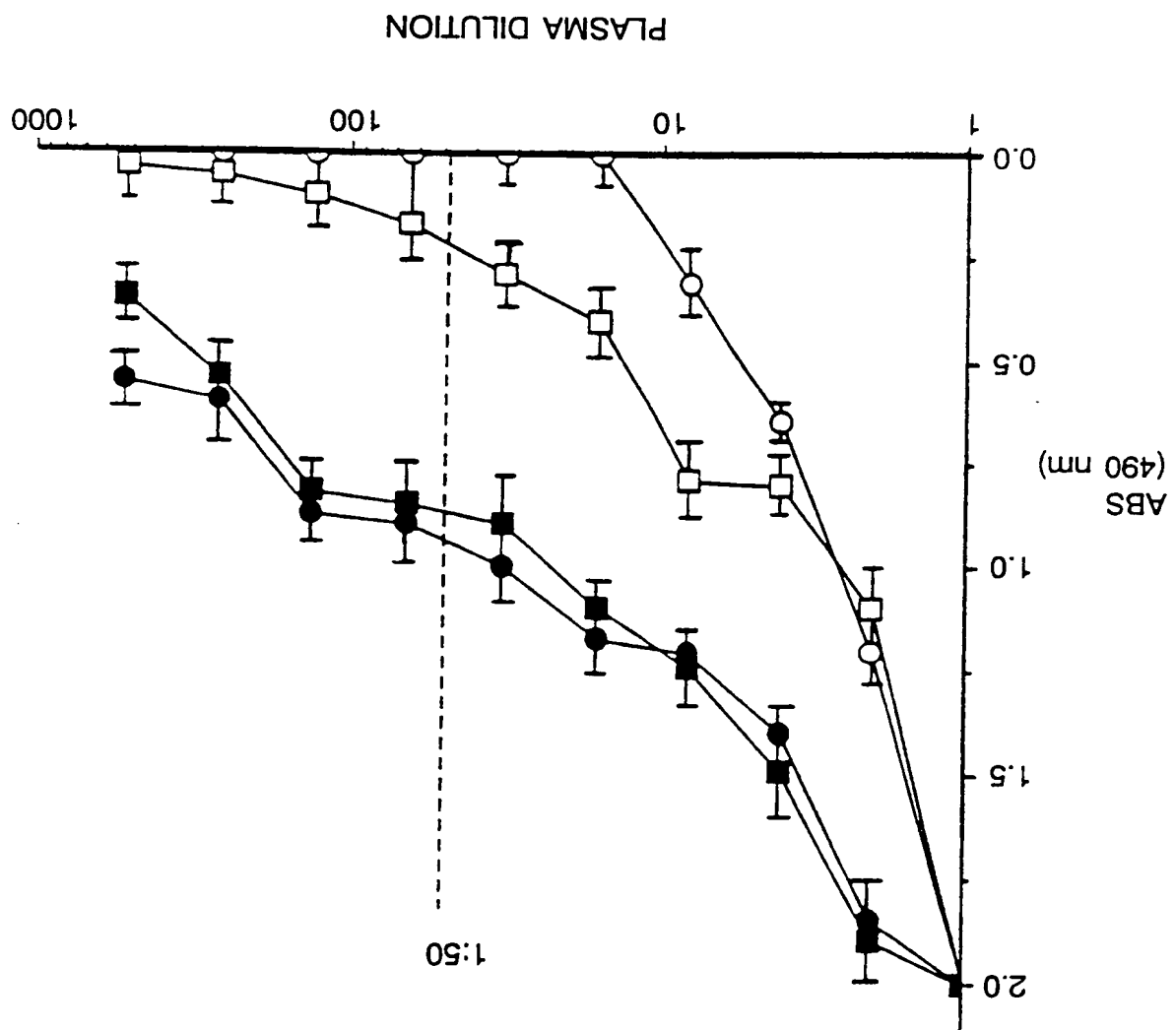
LCAP (U/ml)



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SUSZANNE L. HARRIS

FIG. 8



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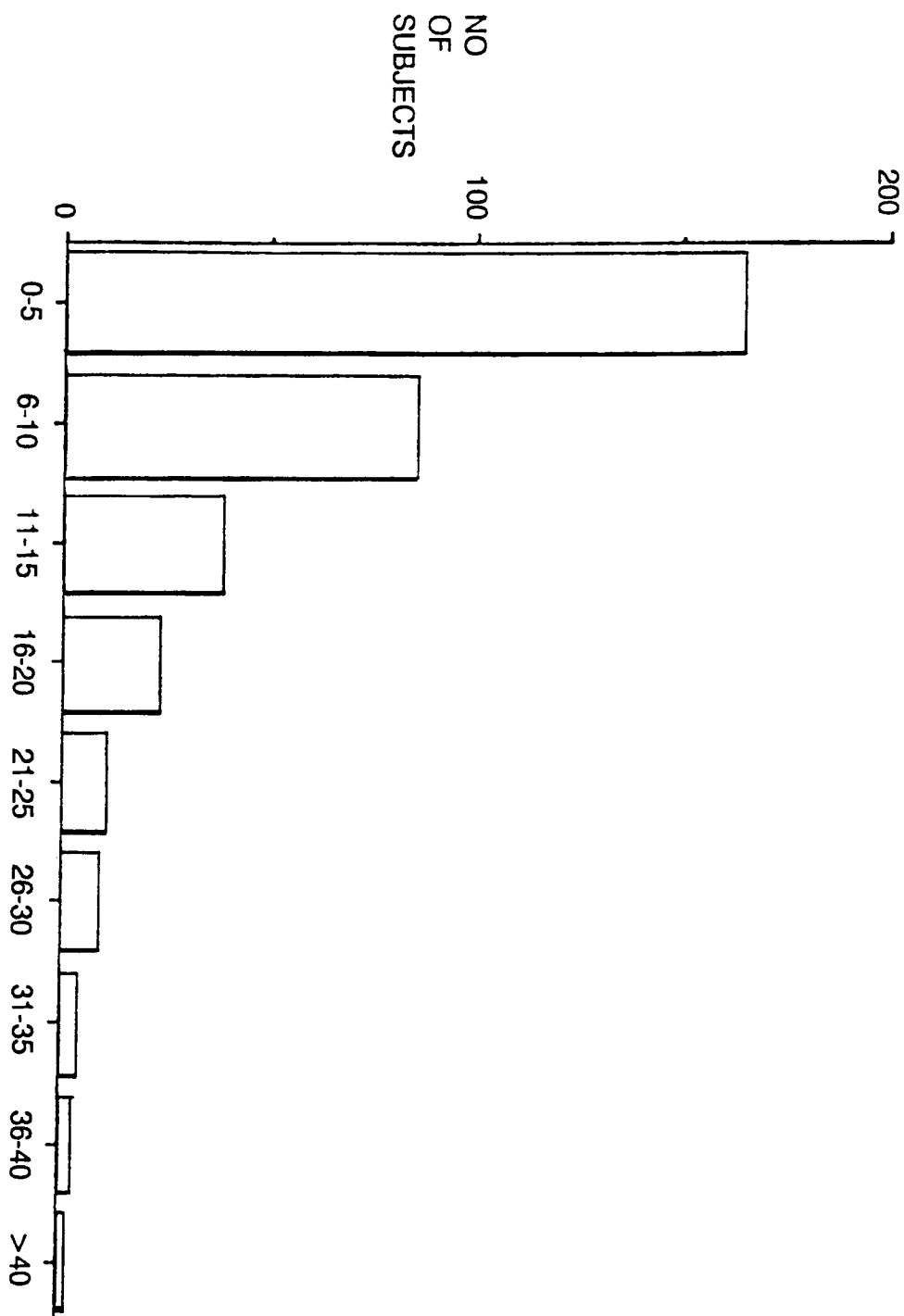


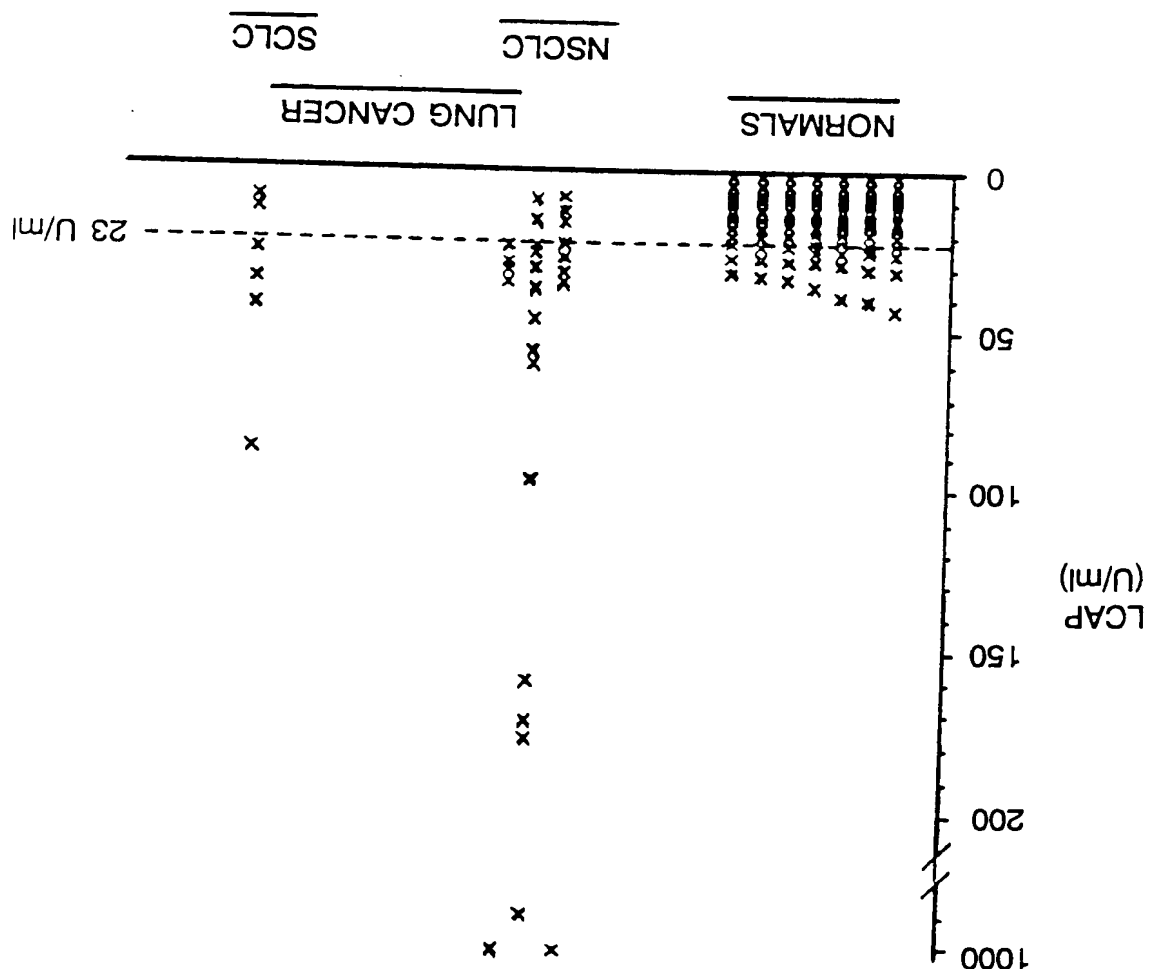
FIG. 9

LCAP (U/ML)

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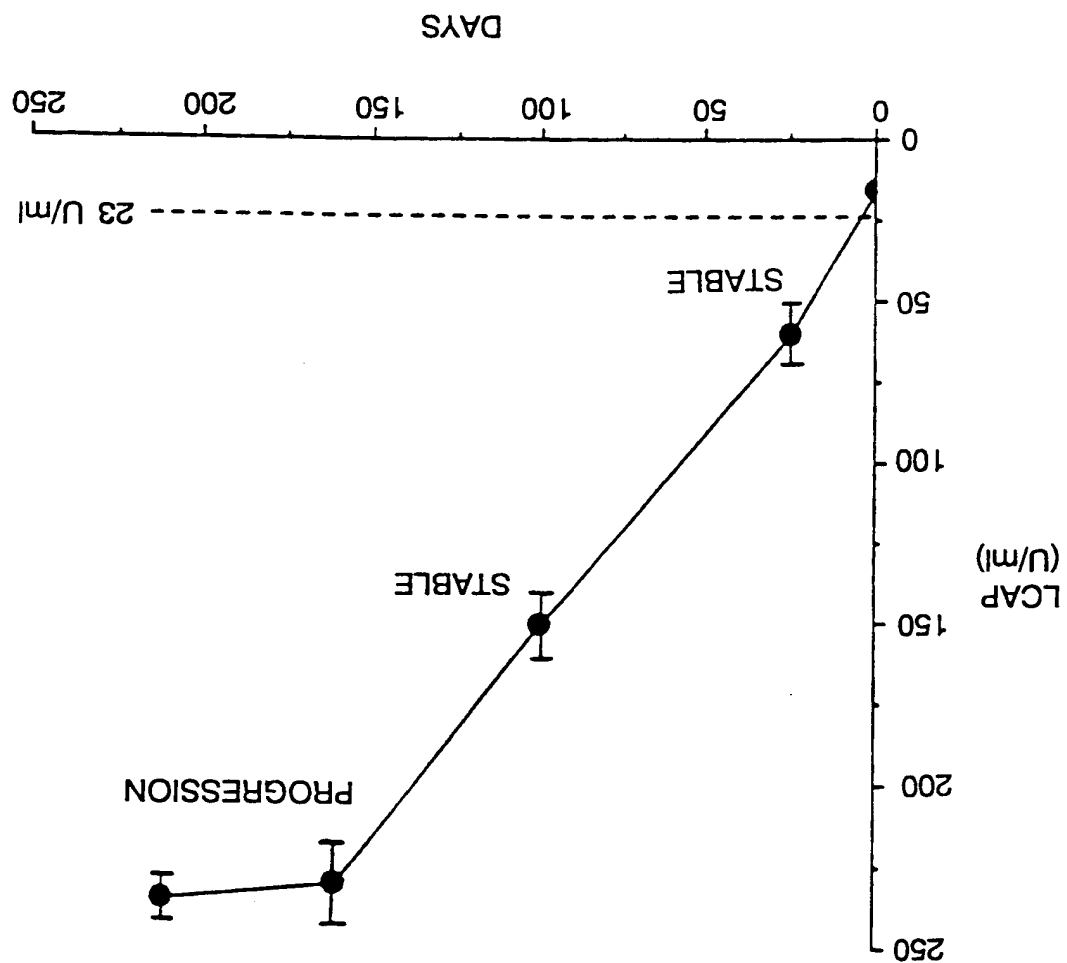
FIG. 10



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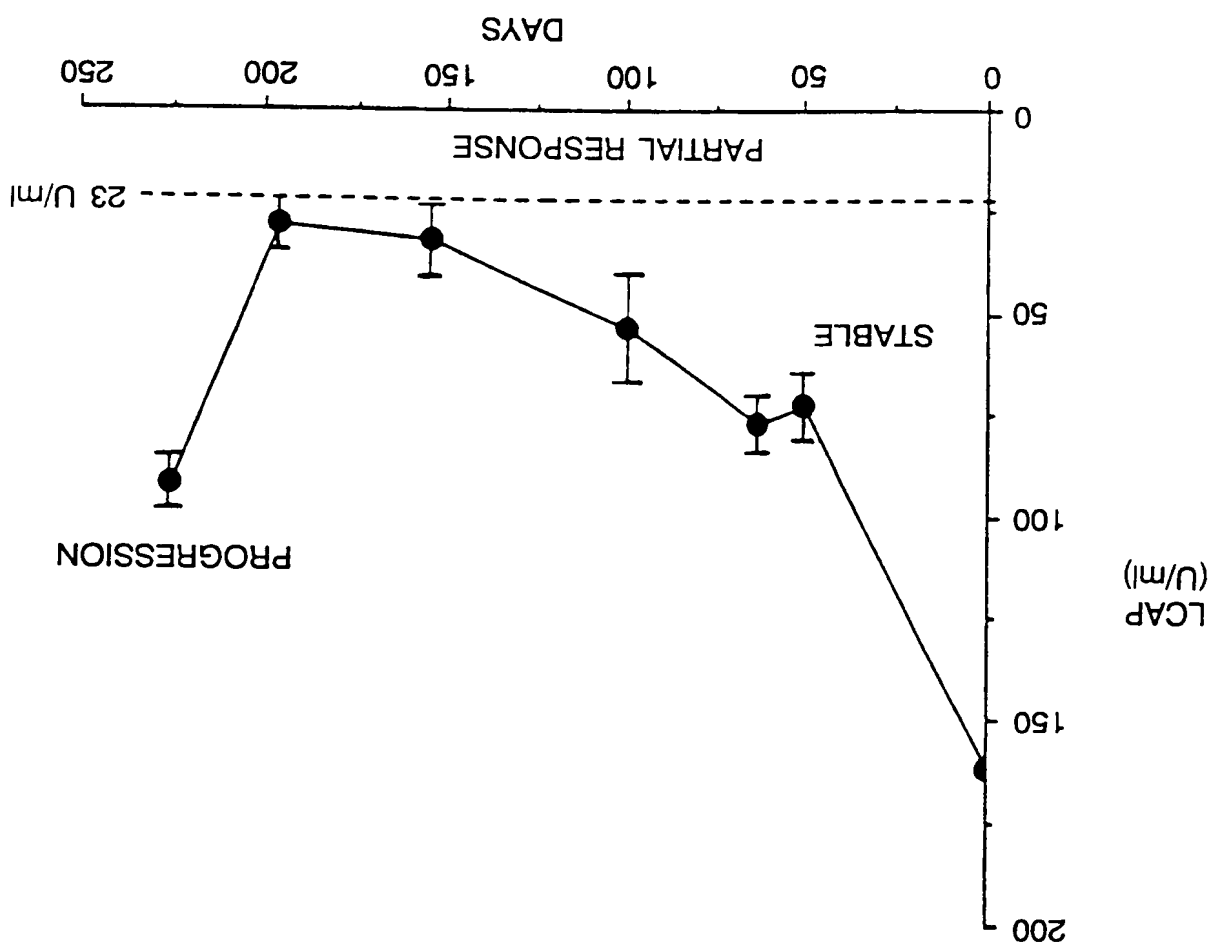
FIG. 11a



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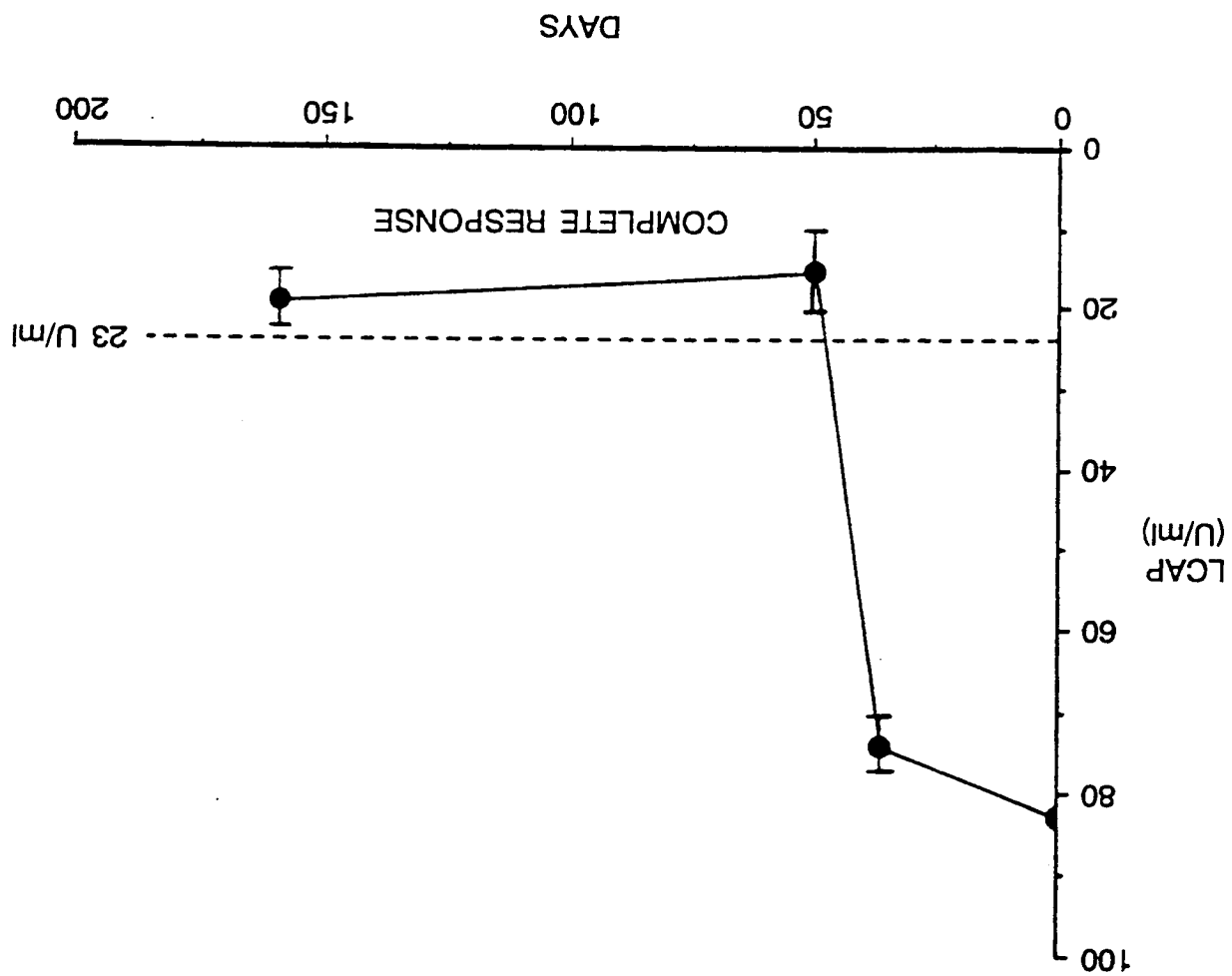
FIG. 11b



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SUBSTITUTED SHEET

FIG. 11C



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SUBSTITUTE SHEET

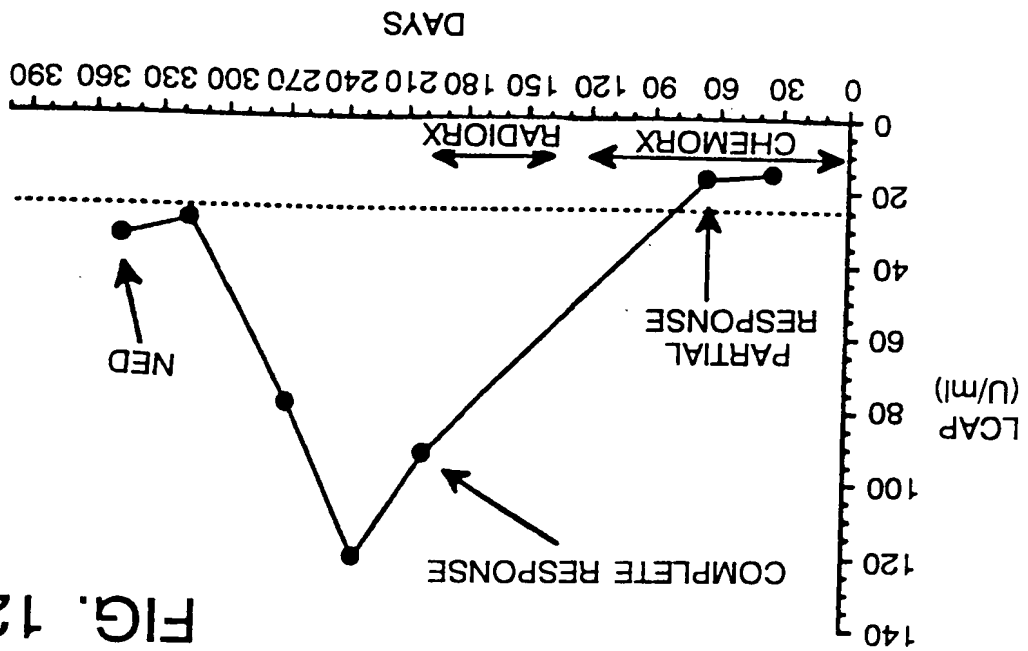


FIG. 12C

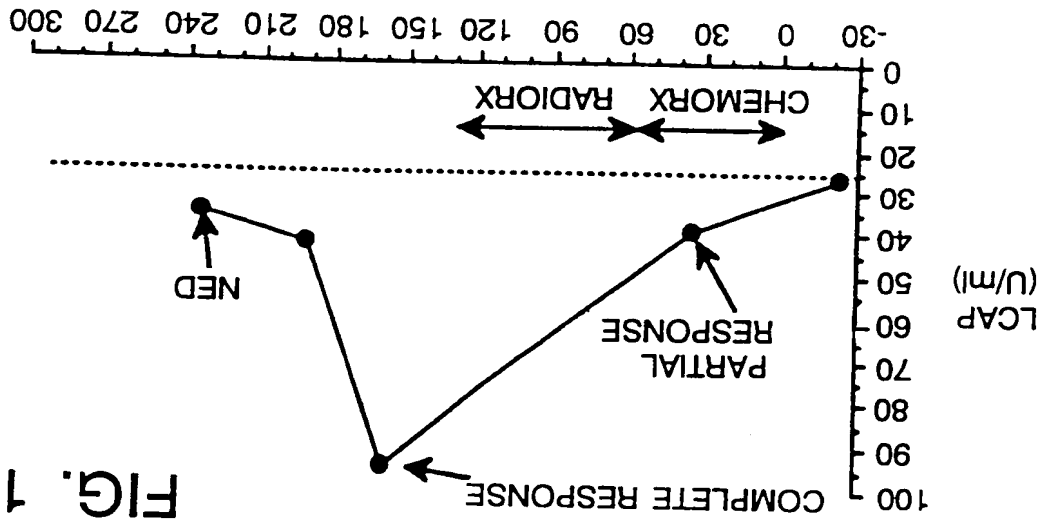


FIG. 12B

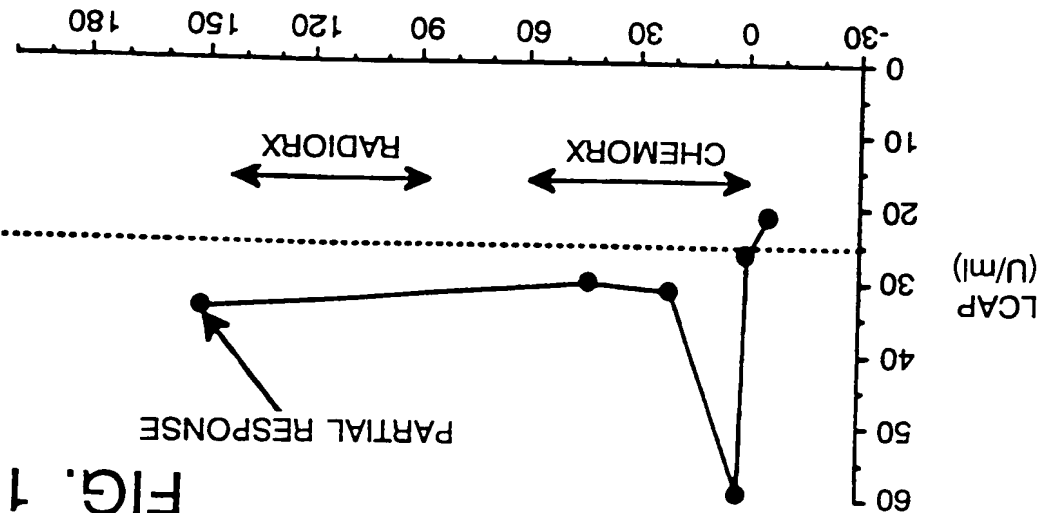


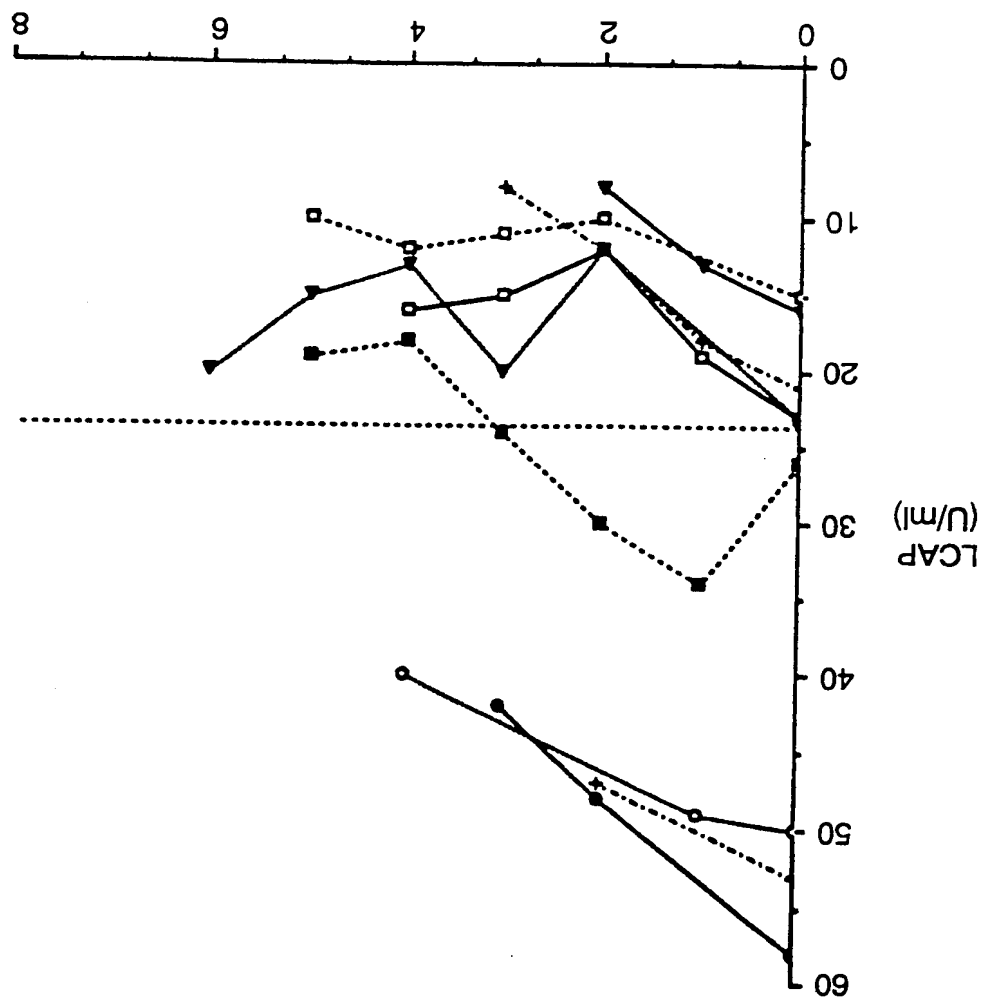
FIG. 12A

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SUBSIDIARY STATEMENT

FIG. 13

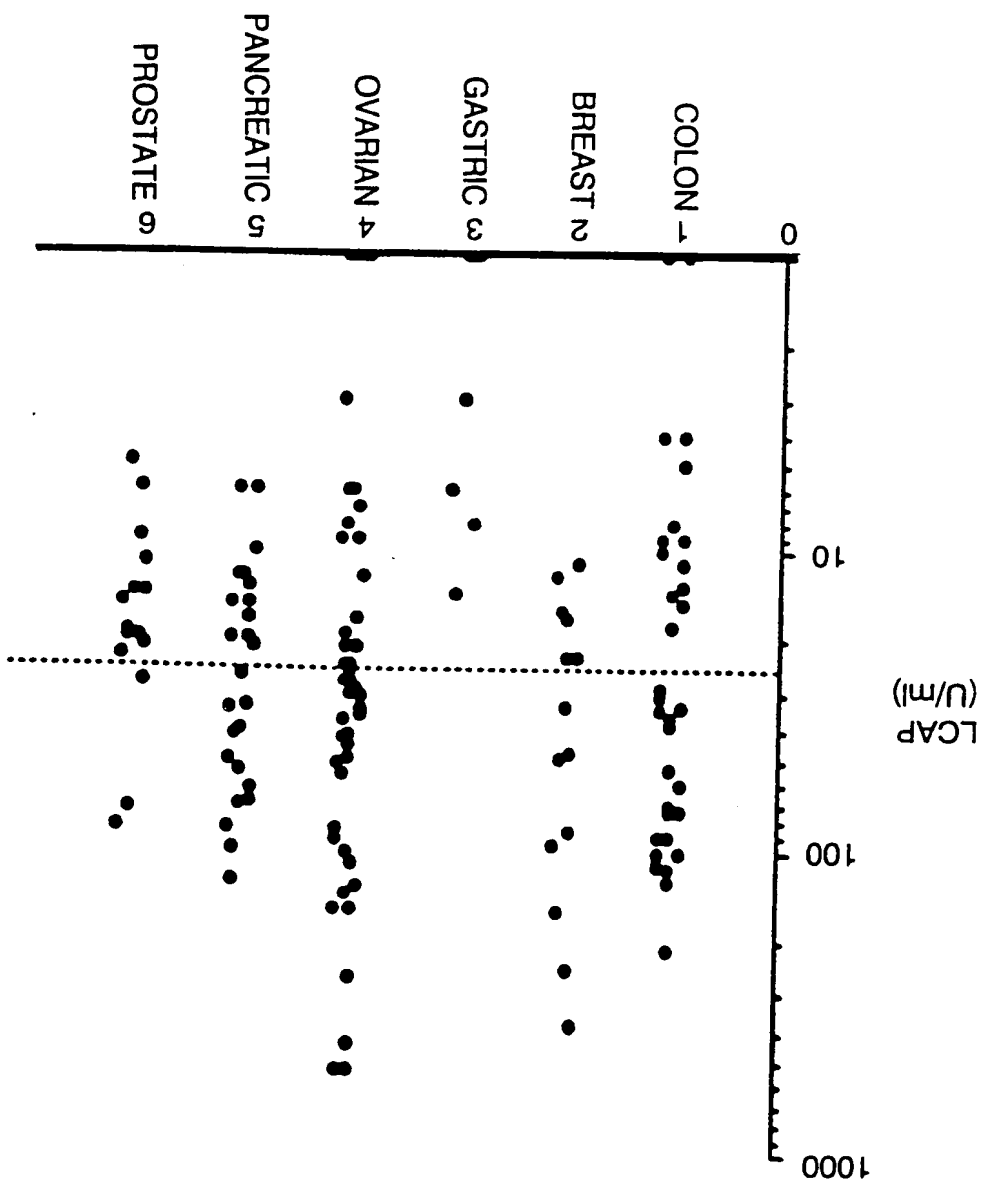
DAYS



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SUBSTITUTION OF LCEP

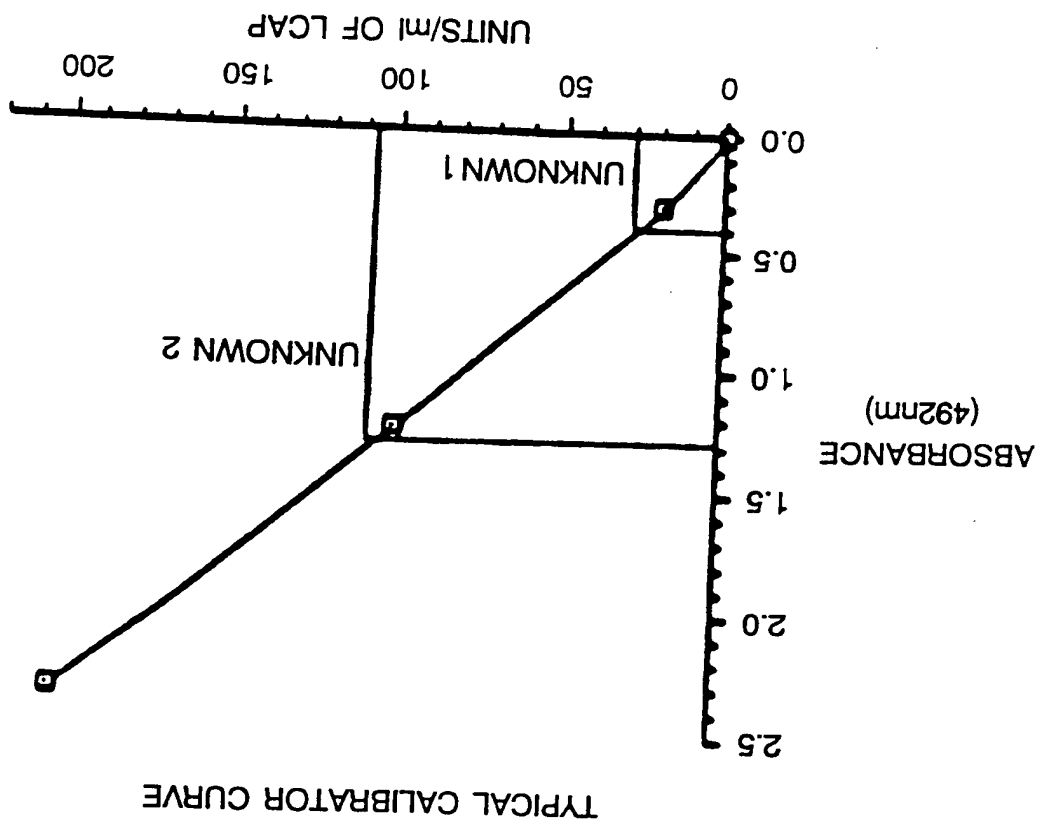
FIG. 14



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SUGAR ANALYSIS

FIG. 15



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U.S. CL. #:		30/300		II. PLEIDS SEARCHED	
According to International Patent Classification (IPC) or to both National Classification and IPC					
CLASSEIFICATION OF SUBJECT MATTER (II) Search Classification Symbols Apply, Markets All II)					
Accredited to International Patent Classification (IPC) or to both National Classification and IPC					
IPC (5): A61K 39/00					
Classification System					
Minimum Documentation Searched?					
Classification Symbols					
30/300; 435/7.1, 7.23, 7.32					
APs					
Categorization					
Citation of Document, if, with indication, where appropriate, of the relevant passages in					
Cancer Research, Volume 44, Issued May 1984, Rosen et al., "Analyses of Human Antigens Using a Panel of Rat Monoclonal Antibodies", pages 2052-2061.					
Cancer Research, Volume 44, Issued November 1984, Okaabe et al., "Monoclonal Antibodies to Surface Antigen of Small Cell Carcinoma of the Lung", pages 5273-5278.					
Cancer, Volume 65, Issued 15 March 1990, Chresten et al., "An Analyses of Immunochemical Excesses for the Detection of the Early Stages of Colon Cancer", pages 1338-1344, see entire document.					
Cancer Research, Volume 50, Issued 15 October 1990, Matmonis et al., "Detection and Characterization of a High Molecular Weight Human Lung Carcino-associated Antigen", pages 673B-6743.					
Cancer Research, Volume 44, see entire document.					
I-10, 43-45					
III. DOCUMENTS CONSIDERED TO BE RELEVANT					
Citation of Document, if, with indication, where appropriate, of the relevant passages in					
Cancer Research, Volume 44, Issued May 1984, Rosen et al., "Analyses of Human Antigens Using a Panel of Rat Monoclonal Antibodies", pages 2052-2061.					
Cancer Research, Volume 44, Issued November 1984, Okaabe et al., "Monoclonal Antibodies to Surface Antigen of Small Cell Carcinoma of the Lung", pages 5273-5278.					
Cancer, Volume 65, Issued 15 March 1990, Chresten et al., "An Analyses of Immunochemical Excesses for the Detection of the Early Stages of Colon Cancer", pages 1338-1344, see entire document.					
Cancer Research, Volume 50, Issued 15 October 1990, Matmonis et al., "Detection and Characterization of a High Molecular Weight Human Lung Carcino-associated Antigen", pages 673B-6743.					
I-10, 43-45					
IV. CERTIFICATIONS					
Signature of Authorized Officer					
Date of Making of this International Search Report					
10 Feb 1992					
International Searching Authority					
Jeffrey Stucker					
Jeffrey Stucker					

- No further action was taken by the patentee or assignee of additional search fees.
- The additional search fees were assessed without filing an additional fee, the Informational Searcher's process.

A full searchable claims could be searched without filing an additional fee, the Informational Searcher's Authority did not require payment of any additional fee.

(Telephone Practice) I-10, 43-45

No unusual additional search fees were filed by claim numbers; it is covered by the application, the Informational Search report is restricted to those claims of the Informational application for which fees were paid, specifically claims:

- As only some of the required claims were filed by the applicant, the Informational search report covers all searchable claims of the Informational application.
- All requested additional fees were filed by the applicant, the Informational search report covers all searchable claims

Please see attached sheet.

The Informational Searcher's Authority found no further information to this Informational application as follows:

VI. OBSERVATIONS WHERE UNIT OF INVENTION IS LACKING:

- Claim numbers _____ because they are dependent claims not cited in association with the second and third derivatives of PCT Rule 6(a).

Claims to such an extent that no meaningful information search can be carried out; specificity:

Claim numbers _____ because they relate to subject matter not required to be searched by this Authority, namely:

This informational search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons:

V. OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND USELESS:

VI. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

PCT/US91/07585

- I. CLA1MS 1-10, & 43-45 drawn to a lung cancer-absorbed protein.
 - II. CLA1ME 11-13 drawn to a cell line which produces LCA1.
 - III. CLA1ME 14-16 drawn to a monoclonal antibody against LCA1.
 - IV. CLA1M 17 drawn to a method of producing a monoclonal.
 - V. CLA1ME 18-23 drawn to a method of detecting LCA1.
 - VI. CLA1MS 24-31 drawn to a Kt1.
 - VII. CLA1MS 32-35 drawn to an immunotoxin.
 - VIII. CLA1MS 36-39 drawn to an imaging agent.
 - IX. CLA1MS 40-42 drawn to a method of detecting tumors.
 - X. CLA1MS I-10, & 43-45 drawn to a lung cancer-absorbed product.
- Line which can produce a protein. Group III is drawn to a monoclonal antibody and group IV is drawn to a method of marking the monoclonal antibody. Group V is a method of marking the monoclonal antibody and group VI is an immunotoxin and group VII is drawn to an antibody agent and group VIII is drawn to a method of detecting tumors. The products of groups I, II, III, IV, and V are separate and distinct methods. The groups IV, V, and IX are separate and distinct methods. The groups VI, VII, and VIII are separate and distinct methods. The groups VII and VIII are linked so as to form a single general inventive concept. PCT Rule 13.2 and 13.2 do not provide for multiple products and methods.